

STIC Search Report Biotech-Chem Library

STIC Database Tracking Number: 171411

TO: Minh-Tam Davis

Location: REM-3A24&3C18

Art Unit: 1642

Wednesday, November 16, 2005 Case Serial Number: 09/856812 From: Barb O'Bryen

Location: Biotech-Chem Library

Remsen 1a69

Phone: 571-272-2518

MOB

barbara.obryen@uspto.gov

Search Notes	



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STIC-Biotech/ChemLib

146411

From:

Davis, Minh-Tam

Sent:

Monday, November 14, 2005 11:47 AM

To:

STIC-Biotech/ChemLib

Subject:

Search request for 09/856812

Please search in commercial database, issued patent files, PGPUB:

1) SEQ ID NO: 42 with and without size limitation for the sequences in the database to the size of the sequence.

2) SEQ ID NO:1

Thank you. MINH TAM DAVIS

ART UNIT 1642, ROOM 3A24, MB 3C18

272-0830

CRFC

Searcher:
Searcher Phone:
Date Searcher Picked up:
Date completed: 11 16 p5
Searcher Prep Time:
Online Time:

Ту	pe of Search
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Encode/T	ransl:
Structure	#:Text:
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endors and cost where applicable
STN:
DIALOG:
QUESTEL/ORBIT:
LEXIS/NEXIS:
SEQUENCE SYSTEM:
WWW/Internet:
Other (Specify)

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GenCore version 5.1.6 (c) 1993 - 2005 Compugen Ltd.
                  Copyright
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OM protein - protein search, using sw model

November 15, 2005, 15:28:51; Search time 233.31 Seconds (without alignments) 611.696 Million cell updates/sec uo:

Run

US-09-856-812B-1 Perfect score:

1891 1 MPRAPKRQRCMPEEDLQSQS.......DTTAMASASSSATGSFSYPE 369 BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table: Sequence:

2105692 seqs, 386760381 residues Searched: Total number of hits satisfying chosen parameters:

Post-processing: Minimum Match 0% Maximum Match 100% Minimum DB seq length: 0 Maximum DB seq length: 2000000000

Listing first 45 summaries A_Geneseq_16Dec04:*

geneseqp2003as:* geneseqp2003bs:* geneseqp1980s:* geneseqp1990s:* geneseqp2001s:* geneseqp2002s:* geneseqp2000s:* Database :

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

geneseqp2004s:*

SUMMARIES

Description	Aay71485 Human MAG	Aab80297 Human pro	Abr48215 Human bla	Abu56516 Lung canc	Abo58424 Human gen	Adr09600 Human pro	Aab08736 Amino aci	Aab08734 Amino aci	Abm82457 Tumour-as	Ada83770 Human MAG	Ado59164 Human MAG	Abu56545 Lung canc	Ado43756 Human MAG	Aab08735 Amino aci	Abu56684 Lung canc	Abu56484 Lung canc	Ada83772 Human MAG	Abu04413 Human exp	Abu04425 Human exp	Abu04421 Human exp	Abu04456 Human exp	Abu04454 Human exp	Abo25122 Human MAG	Adn39048 Cancer/an	Abu04416 Human exp
. aı	AAY71485	AAB80297	ABR48215	ABU56516	ABO58424	ADR09600	AAB08736	AAB08734	ABM82457	ADA83770	ADO59164	ABU56545	ADO43756	AAB08735	ABU56684	ABU56484	ADA83772	ABU04413	ABU04425	ABU04421	ABU04456	ABU04454	AB025122	ADN39048	ABU04416
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& Query Match	100.0	100.0	100.0	100.0	97.9	48.2	48.2	48.1	48.1	45.2	45.2	44.9	44.9	44.8	44.8	44.8	44.8	44.8	44.8	44.8	44.8	44.8	44.8	44.8	44.4
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ALIGNMENTS

AAY71485 standard; protein; 369 AA. RESULT 1 **AAY71485**

AAY71485;

(first entry) 12-OCT-2000

Human MAGE-A10 protein.

MAGE-AlO; human; Tumour Rejection Antigen; TRA; Human Leucocyte Antigen; Haf, Major Histocompatibility Complex; MHC; cytolyvic T-1ymphocyte; CTL; immune response stimulator; prophylaxis; therapy; diagnosis; tumour; cancer; TNP; tumour necrosis factor; cytostatic.

Homo sapiens.

WO200032769-A2

08-JUN-2000.

99WO-IB002018. 26-NOV-1999;

98GB-00026143. 27-NOV-1998; (LUDW-) LUDWIG INST CANCER RES.

De Plaen B, Van Pel A, Brasseur F, Huang L,

Boon T;

WPI; 2000-412317/35. N-PSDB; AAD01311, AAD01312, AAD01313.

Novel polypeptides expressed in tumor cells useful for treating cancers have an ability to complex with a major histocompatibility complex molecule and comprises a specific unbroken amino acid sequence.

Claim 1; Fig 7; 80pp; English.

decappetide sequences, that function as tumour rejection antigens (TRAs). These peptides are capable of forming a complex with major. These peptides are capable of forming a complex with major histocompatibility complex (MRC) molecule type HLA-A2.1 (Human Leucocyte Antigon), that are recognised by T.lymphocytes and elicit an immune response from cytolytic T-lymphocytes (CTL). They function as an immune response stimulator. Tumour rejection antigens are useful in prophylaxis, therapy and diagnosis of tumours and are effective in controlling or preventing tumour growth. The present sequence is the human MAGE-AlO protein, comprising nonapeptides and decapeptides, that serve as tumour patent discloses MAGE-A10 and MAGE-A8 polypeptide, nonapeptide and The

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rejection antigens (TRAs). The novel TRAs encoded by MAGE-Al0 is identified using melanoma cell line (LB1751-MEL), stimulated by autologous CTL clone (447A/5) to produce TNF (tumour necrosis factor). Expression of MAGE-Al0 has been detected in a variety of tumours like melanomas, carcinomas of the head and neck, bladder and prostate, myelomas and lung cancer. The only normal tissue expressing MAGE-Al0 is
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                                                                                                             Length 369;
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                                                                                                          100.0%; Score 1891; DB 3;
100.0%; Pred. No. 4.5e-156;
iive 0; Mismatches 0;
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13-AUG-1999; 99US-01486BLP.
17-AUG-1999; 99US-0149173P.
06-OCT-1999; 99US-0158004P.
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The present invention relates to human secreted prostate cancer antigen coding sequences (AAP72741-AAP72789) and proteins (AAB802731-AAB802731).

The coding sequences and proteins of the present invention are useful for preventing, treating or ameliorating a medical condition; and for the diagnosis and treatment of diseases and disorders. Diseases and disorders can be diagnosed and treated include (auto)immune diseases and disorders can disorders (e.g. asthma), hyperproliferative disorders (e.g. arfat versus host disorders (e.g. asthma), hyperproliferative disorders (e.g. cancers and leukemias), carebrovascular disorders (e.g. stocke), arterial occlusive disorders (e.g. arteriosclerosis), angiogenesis related disorders (e.g. crinopathy and keloid scars), ocular disorders (e.g. glaucoma), neurological disorders (e.g. Alzheimer's, Parkinson's disease, epilepsy and Creutzfeld-Jakob disease) and infections caused by bacteria, fungi, viruses or parasites They may also be useful for wound healing, epithelial cell proliferation, supporting cell culture, tissue regeneration, birth control and as a food additive or preservative
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                                                                                                                                                                                                                                       New nucleic acid molecule encoding human secreted prostate cancer antigens, useful for the diagnosis and treatment of disorders such cancer, leukemia and autoimmune disease.
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100.0%; Pred. No. 4.5e-156;
tive 0; Mismatches 0;
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HUMAN GENOME SCI INC. ROSEN C A.
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Best Local Similarity 100.0
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  13-NOV-2001;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     09-NOV-2001;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                ABU56516;
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                                                                                                                                                                                                                                                                                                                                    RESULT
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           The present invention describes a method for detecting a bladder cancerassociated transcript in a cell from a patient. The method comprises contacting a blological sample from the patient with a polymucleotide that selectively hybridises to a sequence that is 80 % identical to a table of sequences (see ACC50951 to ACC51059). ACC50951 to ACC51059 cencode the human bladder cancer-associated proteins given in ABR48146 to ABR48242). Bladder cancer-associated proteins given in ABR48146 to have cytostatic activities, and can be used in artisense gene therapy and in vaccine production. The method can be used for detecting a bladder cancer-associated transcript in a cell from a patient. The method is useful in diagnosting or treating bladder cancer and in screening for compounds that modulate bladder cancer, such as hormones or antibodies. The nucleic acid molecules from the present invention may be used in various screening and diagnostic methods, and for gene therapy, vaccine and/or antisense/inhibition applications
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Detecting a bladder cancer-associated transcript in a cell from a patient, comprises contacting a biological samplé from the patient with bladder cancer-associated polynucleotide or antibody.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL
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                                                                                 Human bladder cancer associated protein sequence SEQ ID NO:149,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              100.0%; Score 1891; DB 6; Length 369; 100.0%; Pred. No. 4.5e-156;
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                                                                                                                                        Human; bladder cancer; cytostatic; gene therapy; vaccine
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                                                                                                                                                                                                                                                                                                                                                                                                                   03-JUL-2001; 2001US-0302814P.
03-AUG-2001; 2001US-0310099P.
08-NOV-2001; 2001US-0343705P.
12-NOV-2001; 2001US-0350666F.
12-APR-2002; 2002US-0372246P.
                                                                                                                                                                                                                                                                                                                                                                 03-JUL-2002; 2002WO-US021338
                             (first entry)
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Matches 369; Conservative
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N-PSDB; ACC51029.
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                                                                                                                                                                                                                                                MO2003003906-A2.
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                                                                                                                                                                                               Homo sapiens
                               12-JUN-2003
                                                                                                                                                                                                                                                                                                          16-JAN-2003
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The invention relates to a method for detecting a lung cancer-associated transcript in a cell from a patient, comprising contacting a biological sample from the patient with a polymucleotide that selectively hybridises to a sequence that is at least 80 % identical to a gene that exhibits increased or decreased expression in lung cancer samples. Lung cancer. associated polymucleotides and polympetides are used for identifying a compound that modulates a lung cancer-associated polympetide, for inhibiting proliferation of a lung cancer-associated cell to treat lung cancer in a patient and for treating a mammal having lung cancer by administering a modulatory compound identified. The methods are useful for treating lung cancer, such as small cell lung cancer, non-small cell lung cancer or other benign or precancerous lesions, e.g. atelectasis, lung cancer or other benign or precancerous lesions, e.g. atelectasis, chypersensitivity pneumonitis, interstitial pulmonary fibrosis, atthma and bronchiectasis. The genes, polymotaeotides and polympetides are useful cor diagnostic purposes and as targets for screening for therapeutic
                                           360
                                                                                                                                                                                                                                                                                                                                                                                                             Lung cancer-associated polypeptide; cytostatic; emphysema; atelectasis; antinflammatory; antiasthmatic; non-small cell lung cancer; atelectasis; small cell lung cancer; benign lesion; precancerous lesion; bronchitis; chronic obstructive pulmonary disease; hypersensitivity pneumonitis; interstitial pulmonary fibrosis; fibrosis; asthma; bronchiectasis.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Detecting a lung cancer-associated transcript in a cell from a patient for treating lung cancer, by contacting a biological sample from the patient with a polynucleotide that exhibits increased or decreased expression in lung cancer.
                         RAHAEIRKWSLLKFLAKVNGSDPRSFPLWYEEALKDEEERAQDRIATTDDTTAMASASS
RAHABIRKMSLLKFLAKVNGSDPRSFPLWYBBALKDEBBRAQDRIATTDDTTAMASASS
                                                                                                                                                                                                                                                                                                                                                                        Lung cancer-associated polypeptide #109.
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                                                                                                                                                                                                                                           ABUS6516 standard; protein; 369 AA.
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, 2001US-0290492P.
, 2001US-0339245P.
, 2001US-0350666P.
, 2002US-0334370P.
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surveying tissues,
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                                                                                                                                                                                                                                                                                                                                                         New human genome-derived single exon nucleic acid probes useful for human gene expression analysis, for identifying or characterizing alternative splicing events, for assessing genomic alterations or as tools for
                                                                                                                                                                                                                                                                              SSCYPLIPSTPREVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
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compounds that modulate lung cancer, such as antibodies. Sequences ABU56408-ABU56745 represent lung cancer-associated polypeptides of the
                                                                                                                                                                                                  1 MPRAPKRORCMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSSSSSSSS
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                                                                                                                                                             Gaps
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                                                                                                                   Length 369;
                                                                                                                                                           Indels
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                                                                                                                   100.0%; Score 1891; DB 6; 100.0%; Pred. No. 4.5e-156;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human genome derived single exon protein #4658.
                                                                                                                                                           0; Mismatches
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                                                                                                                                                           Matches 369; Conservative
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(RANK/) RANK D R.
(HANZ/) HANZEL D K.
                                                                                                                                        Similarity
                                                                                Sequence 369 AA;
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                                         invention
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                                                                                                                   Query Match
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(HANZ/)
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The invention relates to a nucleic acid probe for measuring human gene expression, comprising any of the 27,400 fully defined nucleotide sequences in the specification, or their complements or fragments, and esquences in the specification, or their complements or fragments, and encoding at least 8 amino acids of any of the 6888 amino acids equences ("Ily defined in the specification. The probe is a single exon probe that daressable set of single exon nucleic acid molecule expression (comprising a plurality of single exon nucleic acid molecule expression (comprising a plurality of single exon nucleic acid molecule expression (comprising a plurality of single exon nucleic acid and addressably solatable or amplifiable from the plurality), a single exon microarray for measuring human gene expression, a vector comprising the single exon and addressably single-exon probe of the plurality of probes is geparately and addressably single-exon probes or microarray for measuring human gene expression, a vector comprising at least 8 contiguous amino acids of any of the above- mentioned amino acid so contiguous amino acids of any of the above- mentioned amino acid so contiguous amino acids of any of the above- mentioned amino acid solated antibody that binds specifically to a peptide cited above, methods of selling and/or licensing single exon probes or microarrays to a customer desiring to measure gene expression, and a computer-readable content record including data on the expression, and a computer-readable content and above. The probe methods and apparatus are useful in gene cypression analysis. The probes may be used as tools for surveying the above. The probes are used in identifying and characterising grounder contain their genemic alterations in the probes are used in identifying and characterising grounder contain their sequence and alterative splicing events, in detecting and characterising grounder and aller genomic alterations in the speciment of the probes of the private of the presence of expression when the pr
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                9
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             RCWITAFGI DVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGYC
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                segdata.uspto.gov/sequence.html?DocID=20030194704
Claim 45; SEQ ID NO 32058; 80pp; English.
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Best Local Similarity 100.0
Matches 362; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Sequence 383 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AT 362
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61 SSCYPLIPSTPERVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120

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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    This invention relates to novel, isolated full length human cDNA molecules and the encoded proteins thereof. Specifically, it refers to cDNA clones obtained by an oligo-capping method, where none of these clones are identical to any known human mRNAs. The present invention describes an immunoassay to identify agonists and antagonists, as well as antiense molecules and siRNAs that can all be used to bind to and modulate expression of the CDNA molecules. As such, these molecules are useful for diagnostic markers or therapeutic targets for the various diseases or morbid states. In particular, they are useful in gene therapy for treating osteoporosis, neurological disease, Alzheimer's disease, Parkinson's disease, dementia, short memory and various cancers, as well as for maintaining equilibrium of sense or motor function, and for treating emotional reaction, fear response and panic. Accordingly, they exhibit osteopathic, neuroprotective, nootropic, antiparkinsonian, cyoteatic and tranquiliser activities. This polypeptide is a protein encoded by a full length human cDNA sequence of the invention. NOTE: This sequence is not given in the sequence listing of the specification but
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 New 1995 cDNA, useful for treating osteoporosis, neurological diseases, Alzheimer's diseases, Parkinson's diseases, dementia and various cancers.
                                                                                                                                                                                     human; oligo-capping method; diagnostic marker; gene therapy; osteoporosis; neurological disease; Alzheimer's disease; Parkinson's disease; dementia; short memory; cancer; sense er motor function; emorional reaction; fear response; panic; osteopathic; neuroprotective; nootropic; antiparkinsonian; cytostatic;
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                                                                                                                                                    Human protein useful for treating neurological disease Seq 3106
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              48.2%; Score 911; DB 8; Length 394;
larity 54.0%; Pred. No. 1.6e-70;
Conservative 53; Mismatches 81; Indel8 ;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Nishikawa T, Iso
Nagai K, Irie R;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Claim 1; SEQ ID NO 3106; 2686pp; English.
                                             ADR09600 standard; protein; 394 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (REAS-) RES ASSOC BIOTECHNOLOGY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                14-FEB-2003; 2003JP-00102207.
09-MAY-2003; 2003JP-00131452.
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                                                                                                                 (first entry)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Isogai T, Yamamoto J,
Wakamatsu A, Ishii S,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              MPI; 2004-583265/57.
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                                                                                                                                                                                                                                                                                                                     Homo sapiens
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                                                                                                                   04-NOV-2004
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
                                                                               ADR09600
         RESULT 6
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The present sequence represents a human MAGE-A11 polypeptide. The specification also describes MAGE-A5, MAGE-A8, MAGE-A9, and MAGE-A11 polypeptides. The MAGE genes encode tumour rejection antigens which complex to human leukocyte antigens (HLAB), and provoke response by autologous, cytolytic T cells. The genes are located in region q28 of the X chromosome. The MAGE polynucleotides are useful for diagnosis of cancer
                                                                                                    240
           173
                                         180
                                                           TPEBVIWEALNAMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYBFLWGP 300
                                                                                                                                                                                 294 IPEEVWWEVLSIMGVYAGREHFLFGEPKRLLTQNWVQEKYLVYRQVPGTDPACYEFLWGP 353
                                                                                                                                                                                                                                                                                                                                                                                                                                                            MAGE-A5; MAGE-A8; MAGE-A9; MAGE-A11; tumour rejection antigen;
human leukocyte antigen; HLA; T cell response; region q28; X chromosome;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Complementary polynucleotide of MAGE family, useful in the diagnosis of
121 QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKABILESVIKNYEDHFPLLFSEAS
                                                                                                    181 ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIFFEGYC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match 48.2%; Score 911; DB 3; Length 429; Best Local Similarity 54.0%; Pred. No. 1.8e-70; Matches 183; Conservative 53; Mismatches 81; Indels 3
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                                                                                                                                                                                                                          RAHABIRKMSLLKFLAKVNGSDPRSFPLWYBBALKDBEB 339
                                                                                                                                                                                                                                            Amino acid sequence of a human MAGE-All polypeptide.
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                                                                                                                                                                                                                                                                                                                                     AAB08736 standard; protein; 429
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      cancer in a patient
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Boon-Falleur T;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       WO200052163-A1
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            120
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                                                                                                                                                                                                                                                         354
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1 MPRAPKRQRCMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSSFPSSFPSSSSSSS 60

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76 MPLEQRSQHCKPEEGLQAQEEDLGLVGAQALQAEEQEAAFFSST------- 119

1 MPRAPKRORCMPEEDLOSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSS 60

Similarity

183;

Best Local Matches 18

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Gарв

22;

Indels

78;

45; Mismatches

Conservative

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LIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQVLPD 125
                                                                                                                                                                                              SESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEASECMLL
                                          KRORCMPEEDLOSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSSSCYP
                                                                188;
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                                                                                                                                                                                                                                                                        186
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    Matches
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                                                          SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
                                                                                                                                                                                                                240
                                                                                                                                                                                                                                  269 VCMQLLFGIDVKEVDPTSHSYVLVTSLNLSYDGIQCNEQSMPKSGLLIIVLGVIFMEGNC 328
                                                                                                                                                                                                                                                                                                              329 IPEEVMWEVLSIMGVYAGREHFLFGEPKRLLTQNWVQEKYLVYRQVFGTDPACYEFLWGP 388
                        154
                                                                                                                                                                                                                                                                                        TPEEVIWEALNAMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP 300
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          MAGE-A5; MAGE-A8; MAGE-A9; MAGE-A11; tumour rejection antigen;
human leukocyte antigen; HLA; T cell response; region q28; X chromosome;
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                                                                              -----INVGTLEELPA-AESPSPPQSPQEESFSPTAMDAIFGSLSDEGSGSQEKEGPSTS
                                                                                                                                                          209 PDLIDPESFSQDILHDKIIDLVHLLLRKYRVKGLITKAEMLGSVIKAYEDYFPEIFREAS
                                                                                                                                    QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS
                                                                                                                                                                                                              ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILLILSIIFIEGYC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Complementary polynucleotide of MAGE family, useful in the diagnosis cancer in a patient.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 318;
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339
                                                                                                                                                                                                                                                                                                                                                                                          of a human MAGE-A8 polypeptide
                                                                                                                                                                                                                                                                                                                                                                    RAHAEIRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEEE
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Pred. No. 1.4e-70;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Example 4; Page 62-64; 72pp; English.
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56.5%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          02-MAR-1999;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                02-JAN-2001
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Best Local Similarity

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104 LESLFREALDEKVAELVRFLLRKYQIKEPVTKAEMLESVIKNYKNHFPDIFSKASBCMQV 163
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                                                                                                                                                                                                                                                                VFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILILSIIFIEGYCTPEEV 245
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            The invention relates to human tumour-associated antigenic target (TAT) polypeptides, and their related nucleic acids. The TAT polypeptides are overexpressed in cancer tissues compared to normal tissues, and may thus serve as effective targets for the diagnosis and treatment of cancer in
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Tumour-associated antigenic target (TAT) polypeptide PRO83395, SEQ:6309
                                                                                                                                                                                                                           IWEALNAMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGPRAHAE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    New tumor-associated antigenic target polypeptides and nucleic acids, useful in preparing a medicament for treating or detecting a proliferative disorder, e.g. breast, lung, colorectal, ovarian or prostate cancer or tumor.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               tumour; diagnosis; cell proliferative disorder; breast cancer; colorectal cancer; lung cancer; ovarian cancer; liver cancer; central nervous system cancer; badder cancer; pancreatic cancer; cervical cancer; melanoma; leukaemia; hybridisation probe; chromosome identification; chromosome mapping; gene mapping;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Tumour-associated antigenic target; TAT; human; overexpression;
                                                                                                                                                                                                                                                                                                                                                           IRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEE 338
                                                                                                                                                                                                                                                                                                                                                                                          ID NO 6309; 7273pp; English,
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABM82457 standard; protein;
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N-PSDB; ACN41095.
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mammals. The invention also relates to nucleic acid and polypeptide sequences at least 80% identical to the TAT nucleic acids and comparising a polypeptides; expression vectors and host cells compirating a TAT nucleic acid; an antibody specific for a TAT polypeptide; a peptide or organic molecule which binds to a TAT polypeptide; fusion proteins comprising a TAT polypeptide; and methods and compositions for the treatment or diagnosis of cancer in mammals. TAT polypeptides, nucleic acids, antibodies, antagonists, binding molecules and compositions are useful for diagnosing or treating a cell proliferative disorder associated with increased TAT expression, particularly cancers such as breast cancer, colorectal cancer, lung cancer, ovarian cancer, liver cancer, bladder cancer, pancreatic cancer, cervical cancer, cancers of the central cancer, melanoma and leukaemia. TAT nucleic acids may further be used as hybridisation probes, in chromosome and gene mapping, in chromosome identification and in gene therapy. The present sequence represents a TAT polypeptide of the invention
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               104 LESLFREALDEKVAELVRFLIRKYQIKEPVTKAEMLESVIKNYKNHFPDIFSKASBCMQV 163
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      245
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   LIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQVLPD 125
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           IWEALNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGPRAHAE 305
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           human; marker; expressed sequence tag; BST; arabidopsis; tumour; stress-induced phenotype; hyperosmotic stress; colon cancer; immunogen;
                                                                                                                                                                                                                                                                                                                                                                                                                               6 KRORCMPEEDLOSOSETOGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSSSSCYP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              SESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEASECMLL
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                                                                                                                                                                                                                                                                                                                                                                                           Gape
                                                                                                                                                                                                                                                                                                                                                                                           22;
                                                                                                                                                                                                                                                                                                                                                       DB 8; Length 318;
                                                                                                                                                                                                                                                                                                                                                       Query Match 48.1%; Score 910; DB 8; Length 316
Best Local Similarity 56.5%; Pred. No. 1.4e-70;
Matches 188; Conservative 45; Mismatches 78; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                  KSORYKAEEGLOAOGEAPGLMDVOIPTAEEOKAASSST
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2001US-0330457P.
2002US-0357144P.
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                                                                                                                                                                                                                                                                                                                     Sequence 318 AA;
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22-OCT-2001; 2
19-FEB-2002; 2
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is a marker for a predetermined phenotype/cell type of interest from a biological species. The method comprises performing a global comparison of a group of expressed sequence tags (ESTS) known to be expressed in the phenotype/cell type of interest with all ESTS expressed in the phenotype/cell type of interest with all ESTS expressed in the in order to identify ESTS that are preferentially expressed in the condition of interest. A method of the invention is useful for determining whether a nucleic acid is a marker for a predetermined phenotype or cell type of interest from a biological species, preferably as a tumour cell, and the predetermined phenotype is a stress-induced phenotype such as hyperosmotic stress or high salt conditions. A method of the invention is also useful for determining the progression of colon cancer in a human, for detecting a tumour cell, and for regulating or preventing the growth of a tumour cell. An antibody of the invention is useful for detecting at the antibody of the invention is useful as an endingent for vaccinating an animal. The present sequence represents a cumour-associated markers. A polypeptide of the invention is useful as an endingent of the invention.
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                                                                                                                                                                                                                                                                                                                                           The invention relates to a novel method for determining if a nucleic acid is a marker for a predetermined phenotype/cell type of interest from a
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    121 QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS 180
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            181 ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGYC 240
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        98
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             44
                                                                                                                                                                        Determining if a nucleic acid is a marker for a phenotype/cell type of interest, by global comparison of expressed sequence tags known to be expressed in the phenotype/cell type with all ESTs expressed in normal
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            --SPDAESLFREALSNKYDELAHFLLRKYRAKELVTKAEMLERVIKOVYKRCFPVIFGKAS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   241 TPEEVIWEALNAMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYBFLWGP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Lobashev AV, Krukovskaya LL;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                1 MPRAPKRORCMPEEDLOSGSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               61 SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  45.2%; Score 854; DB 6; Length 317; ilarity 51.9%; Pred. No. 1.1e-65; Conservative 51; Mismatches 88; Indels 2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   301 RAHABIRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
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                                                                                                                                                                                                                                                                                                Claim 29; Page 156-158; 516pp; English.
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                                                                Baranova AV, Yankovsky NK,
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                     (BIOM-) BIOMEDICAL CENT.
                                                                                                         WPI; 2003-175241/17.
N-PSDB; ADA83769.
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Matches 176; Conserv
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(first entry)

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Lung cancer-associated polypeptide, cytostatic; emphysema; atelectasis; antiinflammatory; antiasthmatic; non-small cell lung cancer; atelectasis; small cell lung cancer; benign lesion; precancerous lesion; bronchitis; chronic obstructive pulmonary diseases; hypersensitivity pneumonitis; interstitial pulmonary fibrosis; fibrosis; asthma; bronchiectasis.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Detecting a lung cancer-associated transcript in a cell from a patient for treating lung cancer, by contacting a biological sample from the patient with a polynucleotide that exhibits increased or decreased expression in lung cancer.
                                             ung cancer-associated polypeptide #138.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Claim 27; Page 294; 453pp; English.
                                                                                                                                                                                                                                                                          ; 2001US-0284770P.
; 2001US-0390492P.
; 2001US-0339245P.
; 2001US-0350666P.
; 2001US-0334370P.
                                                                                                                                                                                                                                                                                                                                                                                (EOSB-) EOS BIOTECHNOLOGY INC
                                                                                                                                                                                                                                                18-APR-2002; 2002WO-US012476
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                                                                                                                                                                                         WO200286443-A2.
                                                                                                                                                                                                                                                                             18-APR-2001;
                                                                                                                                                                                                                                                                                              LO-MAY-2001;
                                                                                                                                                                                                                                                                                                                       13-NOV-2001;
                                                                                                                                                                                                                                                                                                                                                  12-APR-2002;
                                                                                                                                                                                                                                                                                                        09-NOV-2001;
                02-APR-2003
                                                                                                                                                                                                                     31-OCT-2002
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Matches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              --SPDAESLFREALSNKVDELAHFILIRKYRAKELVTKAEMLERVIKNYKRCFPVIFGKAS 156
                                          human; anticancer; melanoma antigen; MAGE-A4; cytostatic; hepatic cancer; gankyrin; Gann ankyrin repeat protein; PSMD10; P28; oncogene;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     61 SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              121 QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS 180
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ECMLLVPGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGYC 240
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | :::|||||||||| ::: ||| |||:||:|| || BSLKMIFGIDVKEVDPTSNTYTLVTCGLSYDGLLGNNQIFPKTGLLIIVLGTIAMEGDS 216
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             TPEEVIWEALANMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP 300
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          217 ASEEEIWEELGVMGVYDGREHTVYGEPRKILITQDWVQENYLEYRQVPGSNPARYEFLWGP 276
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ----PLVPGTLEEVPA-AESAGPPQSPQGASALPTISFTCWRQPNEGSSSQEEEGPSI- 98
                                                                                                                                                                                                                                                                                                                                                                                                    This invention relates to a novel anticancer agent that comprises the melanoma antigen MAGE-A4 related protein. Specifically, it refers to a MAGE-A4 containing Cytostatic agent useful for treating hepatic cancers that highly express gankyrin, where gankyrin is a Gann ankyrin repeat protein also known as PSWD10 or the P28 oncogene. The present invention describes using MAGE-A4 as a preventative or therapeutic agent to suppress gankyrin activity and hence tumourigenesis. This polypeptide sequence is the human MAGE-A4 related protein of the invention.
                                                                                                                                                                                                                                                                                                                    Novel cytostatic agent for cancers expressing gankyrin, comprising melanoma antigen (MAGE)-A4 related protein and having binding property with gankyrin, useful as preventive or therapeutic agent of cancer.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              1 MPRAPKRORCMPREDLOSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSS
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  24;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  45.2%; Score 854; DB 8; Length 317; 51.9%; Pred. No. 1.1e-65; ive 51; Mismatches 88; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       RAHAEIRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
               Human MAGE-A4 related protein SegID 1.
                                                                                                                                                                                                                                                                                                                                                                          Claim 1; SEQ ID NO 1; 22pp; Japanese.
                                                                                                                                                                                      02-OCT-2003; 2003JP-00344979
                                                                                                                                                                                                                  02-OCT-2002; 2002US-0415104P
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity 51.9
Matches 176; Conservative
                                                                                                                                                                                                                                                                            2004-323475/30.
                                                                                                                                                                                                                                             (FUJI/) FUJITA J.
                                                                                                                                                                                                                                                                                         N-PSDB; ADOS9165
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Sequence 317 AA;
                                                                        tumourigenesis.
                                                                                                                              JP2004123752-A.
                                                                                                    gapiens.
                                                                                                                                                           22-APR-2004
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The invention relates to a method for detecting a lung cancer-associated transcript in a cell from a patient, comprising contacting a biological sample from the patient with a polymucleotide that selectively hybridises to a sequence that is at least 80% identical to a gene that exhibits increased or decreased expression in lung cancer associated polymucleotides and polypeptides are used for identifying a compound that modulates a lung cancer-associated polypeptide, for inhibiting proliferation of a lung cancer-associated cell to treat lung cancer in a patient and for treating a mammal having lung cancer by administering a modulatory compound identified. The methods are useful,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ω,
                                                                                                                                                                                                                                                                                                                                                                                                                                 for treating lung cancer, such as small cell lung cancer, non-small cell lung cancer or other benign or precancerous lesions, e.g. atelectasis, emphysema, bronchitis, chronic obstructive pulmonary disease, fibrosis, hypersensitivity pneumonitis, interstitial pulmonary fibrosis, asthma and bronchiectasis. The genes, polymucleotides and polypeptides are useful for diagnostic purposes and as targets for screening for therapeutic compounds that modulate lung cancer, such as antibodies. Sequences there is a such as antibodies. Sequences the such as antibodies are set of the such as antibodies.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            1 MPRAPKRQRCMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 1 MSSEQKSÖHCKPEEGVEAQEEALGLVGAQAPTTEEQEAAVSSSS--------
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 317;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 44.9%; Score 849; DB 6; Length 31 ilarity 51.6%; Pred. No. 3e-65; Conservative 51; Mismatches 89; Indels
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ses 175; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Sequence 317 AA;
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ABUS6545 standard; protein; 317

ABU56545

MPRAPKRORCMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSS

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The present invention relates to MAGE-A4 human leukocyte antigen (HLA) class I-binding peptides, and their use. The antigenic peptides derived from MAGE-A4, when presented by HLA-B37, induce the activation and proliferation of CD8+ cytotocaic T lymphocytes. Also disclosed are the polypeptide and polynucleotide sequences for human MAGE-A4. The peptides of the invention are useful for treating a subject having a disorder with aberrant expression of MAGE-A4. Such disorders particularly include cancer, e.g. metanoma, myeloma or leukaemias. The present sequence
                --SPDAESLFREALSNKVDELAHFLLRKYRAKELVTKAEMLERVIKNYKRCFPVIFGKAS 156
                                                                        | : ::||||||||| ::: ||| |||:||:| | |||| BSLKMIFGIDVKEVDPASNTYTLVTCLGLSYDGLLGNNQIPPKTGLLIIVLGTIAMEGDS 216
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        useful for
QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS
                                                                                                         TPEEVIWEALNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP
                                                                                                                          Human, MAGE-44; human leukocyte antigen; HLA class I-binding peptide;
HLA-B37; CD8+ cytotoxic I lymphocyte; cancer; melanoma; myeloma;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Novel MAGE-A4 human leukocyte antigen class I-binding peptide, u
treating a subject having a disorder with expression of MAGE-A4.
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                                                                                                                                                                                   Boon-Falleur T,
                                                                                                                                                                   RAHABIRKMSLLKFLAKVNGSDPRSFPLWYEBALKDEBE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Claim 4; SEQ ID NO 2; 40pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Russo V,
                                                                                                                                                                                                                                                                  ADO43756 standard; protein; 317
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         13-AUG-2002; 2002US-00218095
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                                                                                                                                                                                                                                                                                                                                                     Human MAGE-A4 polypeptide
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BOON-FALLEUR T.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Zhang Y, Stroobant V,
                                                                                                                                                                                                                                                                                                                                                                                                            leukaemia; cytostatic.
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STROOBANT V.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               2004-447526/42.
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Gaps

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Query Match 44.9%; Score 849; DB 8; Length 317; Best Local Similarity 51.6%; Pred. No. 3e-65; Matches 175; Conservative 51; Mismatches 89; Indels

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The present sequence represents a human MAGE-A9 polypeptide. The specification also describes MAGE-A5, MAGE-A8, MAGE-A9, and MAGE-A11 polypeptides. The MAGE genes encode tumour rejection antispens which complex to human leukocyte antigens (HLA8), and provoke response by autologous, cytolytic T cells. The genes are located in region q28 of the X chromosome. The MAGE polynucleotides are useful for diagnosis of cancer
                                                                                                                                                                                                                                                                                                                                                    300
                                                                             120
                                                                                                                                                                                                   180
                                                                                                                                                                                                                               --SPDABSLFREALSNKVDELAHFLLRKYRAKELVTKAEMLERVIKNYKRCFPVIFGKAS 156
                                                                                                                                                                                                                                                                                                                  240
                                                                                                                                     86
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          MAGE-A5; MAGE-A9; MAGE-A9; MAGE-A11; tumour rejection antigen;
human leukocyte antigen; HLA; T cell response; region q28; X chromosome;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                       217 ASEEEIWEELGVMGVYDGREHTVYGEPRKLLIQDWVQENYLEYRQVPGSNPARYEFLWGP
                                                                                                               ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGYC
                                                                                SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL
                                                                                                                                                                                                   121 QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKABILESVIKNYEDHFPLLFSEAS
                                                                                                                                                                                                                                                                                                                                                                                                                                        241 TPEEVIWEALNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP
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MSSEOKSOHCKPEEGVEAOEEALGLVGAOAPTTEEGEAAVSSS---
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           RAHAEIRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Amino acid sequence of a human MAGE-A9 polypeptide
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RALAETSYVKVLEHVVRVNARVRIAYPSLREAALLEEEE
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                                                                                                     CYPLIPSTPREVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQV 122
                                                                                                                                                                       PAQLEFWFQEALKLKVAELVHFLLHKYRVKEPVTKAEMLESVIKNYKRYFPVIFGKASEF 156
                                                                                                                                                                                                                                                             EEVIWEALNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGPRA 302
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                                                                                                                          ----SKEEEVSAAGSS-SPPQSPQGGASSSISVYYTLWSQPDEGSSSQEEREPSSSVD
                                                                                                                                                                                                                          RAPKRORCMPREDLOSOSETOGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSSS
                                                                            RSP---HCKPDEDLEAQGEDLGLMGAQEPTGEEEETTSSSD-------
                                                                                                                                                        123 LPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEASEC
                                                                                                                                                                                                          MLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGYCTP
                            Gaps
                          29;
    Length 315;
                          Indele
                                                                                                                                                                                                                                                                                                                               HAEIRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
 44.8%; Score 846.5; DB 3;
51.0%; Pred. No. 4.9e-65;
ive 48; Mismatches 88;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  cancer-associated polypeptide #277
                                                                                                                                                                                                                                                                                                                                                                                                       ABU56684 standard; protein; 315 AA
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2001US-0350666P.
2001US-0334370P.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                         (first entry)
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             al Similarity 51.0
172; Conservative
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10-MAY-2001;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          12-APR-2002;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Unidentified
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Query Match
Best Local S:
Matches 172
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Claim 27; Page 401; 453pp; English

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The invention relates to a method for detecting a lung cancer-associated transcript in a cell from a patient, comprising contacting a biological sample from the patient with a polymoleotide that selectively hybridises to a sequence that is at least 80 % identical to a gene that exhibits in creased or decreased expression in lung cancer samples. Lung cancer-associated polymoleotides and polypeptides are used for identifying a compound that modulates a lung cancer-associated polypeptide, for inhibiting proliferation of a lung cancer-associated cell to treat lung cancer in a patient and for treating a mammal having lung cancer in a patient and for treating a mammal having lung cancer to compound identified. The methods are useful for treating lung cancer, such as small cell lung cancer, non-small cell lung cancer or other benign or precancerous lessions, e.g. atclectasis, hypersensitivity pneumonitis, interstitial pulmonary disease, fibrosis, hypersensitivity pneumonitis, interstitial pulmonary fibrosis, asthma and bronchiectasis. The genes, polymucleotides and polypeptides are useful compounds that modulate lung cancer, such as antibodies. Sequences the compounds that modulate lung cancer, such as antibodies. Sequences
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       63 CYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQV 122
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    -----SKEEEVSAAGSS-SPPQSPQGGASSISVYYTLWSQFDEGSSSQEEEEPSSSVD 96
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|SP---HCKPDEDLEAQGEDLGLMGAQEPTGEEEETTSSSD-----
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ||| :::| :| :| :| || || || || HAETSYEKVINYLVMLNAREPICYPSLYEKVLGEEQE 313
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                  Invention
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OM protein - protein search, using sw model

November 15, 2005, 15:32:22 Run on:

; Search time 51.7381 Seconds (without alignments) 686.225 Million cell updates/sec

US-09-856-812B-1 1891 1 MPRAPKRQRCMPEEDLQSQS......DTTAWASASSSATGSFSYPR 369 Title: Perfect score: Sequence:

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

283416 Total number of hits satisfying chosen parameters: 283416 segs, 96216763 residues Searched:

Minimum DB seq length: 0 Maximum DB seq length: 200000000

summaries Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 su

1: pirl:* 2: pir2:* 3: pir3:* 4: pir4:* PIR 79:* Database

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score 1891 911	Query Match 100.0	Len	8 7 7	ID 138659 138660	ના !	en : M M :
12 W 4	911 849 846 5	44 44 2.0.4	319	0 00	138660 138661 138668		
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H84684 T46707	T01532 869703	A45532 T34513 T33152	T4016/ PC4395 T37715	722330 A45546 A48529	A41922 T06609 A84513
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116.5	114.5	114		111.5	110.5
30	333	36 35	3387	544	444

ALIGNMENTS

melanoma antigen MAGE-10 - human C;Species: Homo sapiens (man) C;Date: O'-Jun-1996 #sequence_revision 07-Jun-1996 #text_change 09-Jul-2004 C;Accession: I38659 R;De Plaen, E.; Arden, K.; Traversari, C.; Gaforio, J.J.; Szikora, J.P.; De Smet,

C., Bra

RESULT 1

Immunogenetics 40, 360-369, 1994 A;Title: Structure, chromosomal localization, and expression of 12 genes of the MAGE fam3 A;Reference number: 138659; MUID:95012457; PMID:7927540 A;Accession: 138659

A;Status: preliminary; translated from GB/EMBL/DDBJ
A;Molecule type: DNA
A;Residues: 1-369 «RBS>
A;Crose-references: UNPROT:P43363; EMBL:U10685; NID:g533510; PIDN:AAA68869.1; PID:g53355
C;Genetics:
A;Gene: GDB:MAGEA10; MAGE10

A;Cross-references: GDB:331126
A;Map posttion: Xq28-Xq28
A;Introns: #status absent
C;Superfamily: tumor associated protein MAGB

ö Gape ö Length 369; IndelB Query Match 100.0%; Score 1891; DB 2; Best Local Similarity 100.0%; Pred. No. 9.6e-122; Matches 369; Conservative 0; Mismatches 0;

120 61 SSCYPLIPSTPEBVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKBESPSTL 120 9 SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKERSPSTL 61 g ઠે ઠે

121 QVLPDSBSLPRSBIDBKVTDLVQFLLPKYQMKEPITKABILESVIKAYEDHFPLLFSBAS 180 g ઠે

121 QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS 180 181 ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILISIIFIEGYC 240 ል 8 셤

TPEEVIWEALNMMGLYDGMEHLIYGEPRKILIQDWVQENYLEYRQVPGSDPARYEFLWGP 300 TPEEVIWEALNAMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARXEFLWGP 300 241 181 241 ò g

RAHAEIRKMSLLKFLAKVNGSDPRSFPLWYEBALKDEBBRAQDRIATTDDTTAMASASSS 360 301 301 셤 ሯ

361 ATGSFSYPE 369

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melanoma antigen MAGE-9 - human
C;Species: Homo sapiens (man)
C;Date: 07-Jun-1996 #sequence_revision 07-Jun-1996 #text_change 09-Jul-2004
C;Accession: 138668
R;De Plaen, E.; Arden, K.; Traversari, C.; Gaforio, J.J.; Szikora, J.P.; De Smet, C.; Bracoon, T.
                                                                                                                                                                                                                           A; Experimental source: antigen MAGE-4b
R; Traversari, C.; van der Bruggen, P.; Luescher, I.F.; Lurquin, C.; Chomez, P.; Van Pel,
L; Exp. Mad. 176, 1453-1457, 1992
A; Title: A nonapeptide encoded by human gene MAGE-1 is recognized on HLA-A1 by cytolytic
A; Reference number: PH1294; MUID:93018875; PMID:1402688
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         61 SSCYPLIPSTPREVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILISIIFIEGYC 240
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          A;Residues: 1-172,'T',174-306,'Q',308-317 <DIN>
A;Cross-references: EMBL:U10340; NID:g499123; PIDN:AAA19007.1; PID:g499124
A;Experimental source: melanoma cell line DM150
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   A;Residues: 1-172,'T',174-317 <DEP2>
A;Cross-references: EMBL:U10688; NID:9533516; PIDN:AAA68872.1; PID:9533517
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     A; Molecule type: DNA
A; Residues: 169-172, 'T', 174-177 < TRA2>
A; Experimental source: antigen MAGE-41
A; Darden: Beck, R.J.; Keller, C.J.; Fenton, R.G.
Biochem. Biophys. Res. Commun. 202, 549-555, 1994
A; Title: Cloning and analysis of MAGE-1:related genes.
A; Reference number: JC2358; MUID:94311935; PMID:8037761
A; Accession: JC2359
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            44.9%; Score 849; DB 2;
51.6%; Pred. No. 1.1e-50;
                                                                             A; Status: preliminary; translated from GB/EMBL/DDBJ
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               C;Superfamily: tumor associated protein MAGE P;169-177/Region: HLA-Al binding #status predicted
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             51; Mismatches
source: antigen MAGE-4a
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                A; Experimental source: antigen MAGE-4
A; Accession: PH1298
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                C;Genetics:
A;Gene: GDB:MAGBA4; MAGB4; MAGB-X2
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Matches 175; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                      A; Molecule type: DNA
A; Residues: 169-177 < TRA1>
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A; Introns: #status absent
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                                                                                                                                                                                                                                                                                                                                                                                                                                  A; Accession: PH1297
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A, Reference number: 138659; MUID: 95012457; PMID: 7927540

A, Reference number: 138660

A, Accession: 138660

A, Status: preliminary; translated from GB/EMBL/DDBJ

A, Residues: 1-319 < RES>
A, Residues: 1-319 < RES>
A, Cross-references: UNIPROT: P43364; EMBL: U10686; NID: 9533512; PIDN: AAA68870.1; PID: 95335

C, Genetics:
A, Gene: GDB: MAGEAll; MAGEIl

A, Map position: Xq28-Xq28

A, Introns: #status absent
C, Superfamily: tumor associated protein MAGE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  far
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A;Cross-references: UNIPROT:P43358; EMBL:U10687; NID:g533514; PIDN:AAA68871.1; PID:g5335
                                                                                                                                                                                                                                                                                                                                                 Accession: I38660
De Plaen, B.; Arden, K.; Traversari, C.; Gaforio, J.J.; Szikora, J.P.; De Smet, C.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ..
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            RESULT 3
138661
melanoma antigen MAGE-4 - human
melanoma antigen MAGE-X2
N/Alternate names: MAGE 41 protein; melanoma antigen MAGE-X2
C;Species: Homo sapiens (man)
C;Species: Homo sapien
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ö
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKABILESVIKNYBDHFPLLFSEAS 180
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   99 PDLIDPESFSQDILHDKIIDLVHLLLRKYRVKGLITKAEMLGSVIKNYEDYFPEIFREAS 158
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILISIIFIEGYC 240
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               TPEEVIWEALNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP 300
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ----LINVGTLEELPA-AESPSPPQSPQEESFSPTAMDAIFGSLSDEGSGSQEKEGPSTS 98
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         6
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                                                                                                                                                                                                                                                               Species: Homo sapiens (man)
Date: 07-Jun-1996 #sequence_revision 07-Jun-1996 #text_change 09-Jul-2004
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           219 IPEEVMWEVLSIMGVYAGREHFLFGEPKRLLTQNWVQEKYLVYRQVFGTDPACYEFLWGP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     1 MPRAPKRORCMPEEDLOSOSETOGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               1 MPLEQRSÓHCKPEBGLÓAQEBDLGLVGAQALQAEBQEAAFPSST-------
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       / Match 48.2%; Score 911; DB 2; Length 31 Local Similarity 54.0%; Pred. No. 6.2e-55; Length 31 he 183; Conservative 53; Mismatches 81; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         RAHAEIRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  A;Accession: 138661
A;Status: preliminary; translated from GB/EMBL/DDBJ
A;Molecule type: DNA
A;Residues: 1-317 <DEP1>
                                                                                                                                                                                                                               melanoma antigen MAGE-11 - human
C;Species: Homo aspiens (man)
C;Date: 07-Jun-1996 #sequence_revis
C;Accession: 138660
R;De Plaen, E.; Arden, K.; Traverse
Oon, T.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Immunogenetics 40, 360-369, 1994
||||||||||
ATGSFSYPE 369
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N, Alternate names: tumor-associated antigen, MAGE-3b
   49.3%; Pred. No. 2.1e-47;
                              54; Mismatches
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submitted to the EMBL Data Library, June 1994
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Cross-references: UNIPROT:P43360
Best Local Similarity 49.3% Matches 167; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         melanoma antigen MAGE-6 - human
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A; Residues: 1-314 <DIN>
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                       ArTitle: Structure, chromosomal localization, and expression of 12 genes of the MAGE fam ArReference number: 138659; MUID:95012457; PMID:7927540
A;Reference number: 138668
A;Accession: 138668
A;Status: preliminary; translated from GB/EMBL/DDBJ
A;Molecule type: DNA
A;Residues: 1-315 <RES>
A;Cross-references: UNIPROT:P43362; EMBL:U10694; NID:9533527; PIDN:AAA68877.1; PID:95335
C;Genetics:
A;Gene: GDB:MAGEA9; MAGE9
A;Gross-references: GDB:331125
A;Gross-references: GDB:331125
A;Gross-references: GDB:331125
A;Antons: #status absent
C;Superfamily: tumor associated protein MAGE
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A;Molecule type: DNA
A;Molecule type: DNA
A;Molecule type: DNA
A;Molecule type: DNA
A;Cross-references: UNIPROT: P43356; GB:L18920; NID:9436180; PIDN:AAA17729.1; PID:9436181
B;Traversari, C.; van der Bruggen, P.; Luescher, I.F.; Lurquin, C.; Chomez, P.; Van Pel, J. Exp. Med. 176, 1453-1457, 1992
A;Title: A nonapoptide encoded by human gene MAGE-1 is recognized on HLA-A1 by cytolytic A;Reference number: PH1294; MUID:93018875; PMID:1402688
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   C;Species: Homo sapiens (man)
C;Species: Homo sapiens (man)
C;Species: Homo sapiens
C;Date: 07-Jun-1996 #sequence_revision 07-Jun-1996 #text_change 09-Jul-2004
C;Date: 07-Jun-1996 #sequence_revision 07-Jun-1996 #text_change 09-Jul-2004
C;Accession: 168889; PH1294
F;Pilcs Sequence and expression pattern of the human MAGE2 gene.
A;Reference number: 154519; MUID:94102805; PMID:8276455
A;Accession: 168889
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                97 PAOLEFMFQEALKLKVAELVHFLIHKYRVKEPVTKAEMLESVIKNYKRYFPVIFGKASEF 156
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            63 CYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQV 122
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              123 LPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEASEC 182
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      BEVIWEALNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYBFLWGPRA 302
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ----SKREEVSAAGSS-SPPQSGASSSISVYYTLWSQPDEGSSSQREEPSSSVD 96
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   3 RAPKRORCMPEEDLOSOSETOGLEGAOAPLAVEEDASSSTSTSSSFPSSFPSSSSSSSSS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              MLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILLILSIIFIEGYCTP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 315;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
44.8%; Score 846.5; DB 2; Length
Best Local Similarity 51.0%; Pred. No. 1.6e-50;
Matches 172; Conservative 48; Mismatches 88; Indels
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HAETSYEKVINYLVMLNAREPICYPSLYEBVLGEEQE 313
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   Immunogenetics 40, 360-369, 1994
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A,Residues: 168-176 <TRA>
C,Genetics:
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Immunogenetics 40, 360-369, 1994
A;Title: Structure, chromosomal localization, and expression of 12 genes of the MAGE fam#
A;Reference number: 138659; MUID:95012457; PMID:7927540
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A;Title: A nonspeptide encoded by human gene MAGE-1 is recognized on HLA-A1 by cytolytic A;Reference number: PH1294; MUID:93018875; PMID:1402688
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                                                                                                                                                                                                                                                                                                                                                                      300
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                                                                                                                                                                                             61 SSCYPLIPSTPREVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
                                                                                                                                  44
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Cibecies: Homo sapiens (man)
Cibece: 20-Feb-1995 #sequence revision 20-Feb-1995 #text_change 09-Jul-2004
Cibeces aton: JC2360; PH1301; Is8665; G01445
Ribing, M.; Beck, R.J.; Keller, C.J.; Fenton, R.G.
Blochem. Blochem. Blochem. 202, 549-555, 1994
A;Title: Cloning and analysis of MAGE-1-related genes.
A;Reference number: JC2358; MUID:94311935; PMID:8037761
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A;Molccule type: DN:
A;Residuses: 166-176 <TRA>
R;De Plaen, E.; Arden, K.; Traversari, C.; Gaforio, J.J.; Szikora, J.P.; De
                                                                 1 MPRAPKRORCMPEEDLQSQSETQGLEGAQAPLAVEEDASSSISISSSFPSSSSSSSS
                                                                                                                                                                                                                                                                                                                                   121 QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   181 ECMLLVFGIDVKEVDPTGHSPVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGYC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  241 TPEEVIWEALNWMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP
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A; Molecule type: DNA
A; Residues: 1-314 <RES>
A; Cross-references: EMBL:U10691; NID:g533522; PIDN:AAA68875.1; PID:g533523
                                                                                                                                                                                                                                      A; Cross-references: EMBL: U10339; NID: 9499121; PIDN: AAA19006.1; PID: 9499122
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93,
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A;Molecule type: mRNA
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  A; Map position: Xq28-Xq28
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C.; Bre

DB 2; Length 314;

42.3%; Score 800.5;

Query Match

Superfamily: tumor associated protein MAGE

A;Cross-references: GDB:273684

A; Map position: Xq28-Xq28 A; Gene: GDB: MAGEA2; MAGE2

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A,Gene: GDB:MAGEA12; MAGE12; MAGE-12f
A,Cross-references: GDB:331129
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          A; Experimental source: MAGE-21
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A;Regidues: 168-176 <TRA>
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Best Local Simi
Matches 163;
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A,Experimental source: melanoma cell line DM150
R,Traversari, C.; van der Bruggen, P.; Luescher, I.F.; Lurquin, C.; Chomez, P.; Van Pel, J. Exp. Med. 176, 1453-1457, 1992
J. Exp. Med. 176, 1453-1457, 1992
A,Title: A nonapeptide encoded by human gene MAGE-1 is recognized on HLA-Al by cytolytid A;Reference number: PH1294; MUID:93018875; PMID:1402688
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A; Residues: 168-176 < TRA>
A; Residues: 168-176 < TRA>
B; Gaugler, B; Van den Bynde, B; Van der Bruggen, P.; Romero, P.; Gaforio, J.J.; De Pl.
J. Exp. Med. 179, 921-930, 1994
A; Title: Human gene MAGE-3 codes for an antigen recognized on a melanoma by autologous A; Reference number: I38438; MUID:94157413; PMID:8113684
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: DNA
A; Residues: 1-314 < RES>
A; Cross-references: EMBL:U03735; NID:9468825; PIDN:AAA17446.1; PID:9468826
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                                                                                                                                                                                                                                                                           61 SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPL-DQSDEGSSSQKEESPST 119
                                                                                                                                                                                                                                                                                                                                                                     120 LQVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEA 179
                                                                                                                                                                                                                                                                                                                                                                                               ---PPDLESEFQAALSRKVAKLVHFLLLKYRAREPVTKAEMLGSVVGWQYFFPVIFSKA 154
                                                                                                                                                                                                                                                                                                                                                                                                                                                             SECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGY 239
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     CTPEEVIWEALNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWG 299
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            1 MPRAPKRORCMPEEDLOSOSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSS 60
                                                                                                                                                                                                                                 ----- 44
                                                                                                                                                                                                                                                                                                   Melanoma antigen MAGE-3 - human
MyAlternate names: MAGE 3 protein
Gispecies: Homo sapiens (man)
Gispecies: Homo sapiens (man)
Gibate: 20-Feb-1995 #sequence revision 20-Feb-1995 #text_change 09-Jul-2004
Bibiophys: Res. Commun. 202, 549-555, 1994
A;Title: Cloning and analysis of MAGE-1-related genes.
A;Reference number: JC2258; MUID:94311935; PMID:8037761
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                                                                                          42.2%; Score 798.5; DB 2; Length llarity 49.7%; Pred. No. 2.9e-47; Conservative 58; Mismatches 86; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Indels
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41.4%; Score 783.5; DB 2;
Best Local Similarity 48.5%; Pred. No. 3.1e-46;
Matches 165; Conservative 59; Mismatches 89;
A;Introns: #status absent
C;Superfamily: tumor associated protein MAGE
F;168-176/Region: HLA-Al binding #status predicted
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A,Gene: MAGE-3
C,Superfamily: tumor associated protein MAGE
F,168-176/Region: HLA-Al binding #status predicted
                                                                                                                 Similarity
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A, Molecule type: mRNA
A, Residues: 1-314 <DIN>
                                                                                       Query Match
Best Local Simi
Matches 169;
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A;Cross-references: UNIPROT:P43365; GB:L18877; NID:g499345; PIDN:AAA19023.1; PID:g499346 R;Ding, M.; Beck, R.J.; Keller, C.J.; Fenton, R.G. Biochem. Blophya: Res. Commun. 202, 549-555, 1994 A;Title: Gloning and analysis of MAGE1-related genes.
A;Title: Cloning and analysis of MAGE1-related genes.
A;Reference number: JC2358; MUID:94311935; PMID:8037761
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A; Residues: 1-9, 'S',11-186,'D',188-299,'S',301-314 <DIN>
A; Experimental Bource: melanoma cell line DM150; MAGE-12f
R; Traversari, C.; van der Bruggen, P.; Luescher, I.F.; Lurquin, C.; Chomez, P.; Van Pel,
J. Exp. Med. 176, 1453-1457, 1992
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       melanoma antigen MAGE-12 - human
NyAlternate names: MAGE 21 protein
C;Species: Homo sapiens (man)
C;Date: 07-Jun-1996 #sequence revision 07-Jun-1996 #text_change 09-Jul-2004
C;Date: 07-Jun-1996 #sequence revision 07-Jun-1996 #text_change 09-Jul-2004
C;Date: 07-Jun-1996 #sequence revision 07-Jun-1996 #text_change 09-Jul-2004
C;Accession: 154519; Jun-1295 #Jul-2004
A;Pille: Sequence and expression pattern of the human MAGE2 gene.
A;Reference number: 154519; MUID:94102805; PMID:8276455
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                                                                                                                                                                                                                                                                                                                    ---FPDLESERQAALSRKVABLVHFLLLKYRAREPVTKAEMLGSVVGNWQYFFPVIFSKA 154
                                                                                                                                                                                                                                                                                                                                                                                                                      239
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MPRAPKRORCMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSS
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                                           1 MPLEQRSQHCKPEEGLEARGEALGLVGAQAPATEEQEAASSSST----
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A,Reference number: PH1294; MUID:93018875; PMID:1402688
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ilarity 48.1%; Pred. No. 7.9e-46;
Conservative 55; Mismatches 96;
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A;Molecule type: DNA
A;Residues: 1-314 <DES>
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C,Superfamily: tumor associated protein MAGE
F,168-176/Region: HLA-Al binding #status predicted
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Species: Homo sapiens (man)
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                                                      121 QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS 180
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C.Species: Homo sapiens (man)
C.Species: Nomo sapiens (man)
R.Species: Nomo sapiens (man)
R.Species: Sea. Commun. 202, 549-555, 1994
A.Title: Cloning and analysis of MAGE-1-related genes
A.Reference number: JC2358; MUID:94311935; PMID:8037761
A.Mocession: JC2358
A.Molecule type: mRNA
A.Residues: 1-280 cDIN>
A.Residues: 1-280 cDIN>
C.Genetics:
                                                                    --FPDLETSFQVALSRKWAELVHFLLLKYRAREPFTKAEMLGSVIRNFQDFFPVIFSKAS
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              Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match 40.5%; Score 766; DB 2; Length 280
Best Local Similarity 55.1%; Pred. No. 4.2e-45;
Matches 167; Conservative 34; Mismatches 70; Indels
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C;Superfamily: tumor associated protein MAGE
F;161-169/Region: HLA-Al binding #status predicted
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melanoma antigen MAGE-1 - human
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138008 melanoma antigen MAGE-B1 - human N,Alternate names: MAGE-Xp protein; MAGE-like protein

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Cidence of the Constitution of the Constitutio
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A;Residues: 1.234 <RES>
A;Cross-references: UNIPROT:P43361; EMBL:U10693; NID:g533525; PIDN:AAA68876.1; PID:g53352
C;Genetics:
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R;De Plaen, E.; Arden, K.; Traversari, C.; Gaforio, J.J.; Szikora, J.P.; De Smet,
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        26
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              57 -----IPQKPQ-----GAPPTTTAAAVSCT------ESDEGAKCQGEENASFS
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C;Date: 07-Jun-1996 #sequence_revision 07-Jun-1996 #text_change 09-Jul-2004
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    1 MPRAPKRORCMPEEDLOSOSETOGLEGAQAPLAVEEDASSSTSTSSSFPSSSSSSSS
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A;Title: Structure, chromosomal localization, and expression of 12
A;Reference number: 138659; MUID:95012457; PMID:7927540
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A;Status: prellminary; translated from GB/EMBL/DDBJ
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A,Cross-references: GDB:33123
A,Map position: XQ28-XQ28
A,Introns: #status absenct
C,Superfamily: tumor associated protein MAGE
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hypothetical protein DKPZp434N0522.1 - human (fragment)
C;Species: Homo sapiens (man)
C;Species: Homo sapiens (man)
C;Species: Homo sapiens (man)
C;Accession: T43464
R;Blum, H.; Bauersachs, S.; Mewes, H.W.; Gassenhuber, J.; Wiemann, S.
Bibmithet to the Protein Sequence Database, December 1999
A;Reference number: 222517
A;Accession: T43464
A;Status: preliminary
A;Molecule type: mRNA
A;Residues: 1-476 cAAA>
A;Cross-references: UNIPROT:Q9YSV3; EMBL:AL133628
A;Cross-references: UNIPROT:Q9YSV3; EMBL:AL133628
A;Experimental source: adult testis; clone DKFZp434N0522
C;Genetics:
A;Note: DKFZp434N0522.1
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                                                                                                                                                                             45 LIMGTLEEVT-DSGSPSPPQSPEGASSSLTVTDSTLWSQSDEGSSSNEEEGPSTSPDPAH 103
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                                                                                                                                                                                                                                                    186 VFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILLILSIIFIEGYCTPEEV 245
                                                                                                                                                                                                                                                                                                                                68 PSTPREVSADDETPNPPOSAQIACSSPSVVASLPLDQSDEGSSSQKBESPSTLQVLPDSE 127
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                                                                                                                 Gaps
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 Length 234;
Query Match 33.8%; Score 639; DB 2; Length 23. Best Local Similarity 55.0%; Pred. No. 1.5e-36; Matches 137; Conservative 33; Mismatches 57; Indels
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SGPWSWDD 407
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C;Species: Mus musculus (house mouse)
C;Date: 31-Dec-1991 #sequence_revision 31-Dec-1991 #text_change 09-Jul-2004
C;Accession: JN0148
R;Maruyaman, K.; Usami, M.; Aizawa, T.; Yoshikawa, K.
Biochem. Biophys. Res. Commun. 178, 291-296, 1991
A;Title: A novel brain-specific mRNA encoding nuclear protein (necdin) expressed in neural A;Reference number: JN0148; MUID:91298962; PMID:2069569
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C;Date: 07-Jun-1996 #sequence revision 18-Feb-2000 #text_change 09-Jul-2004
C;Accession: 138663; 138664; PH1299; PH1300
R;De Plaen, E.; Arden, K.; Traversari, C.; Gaforio, J.J.; Szikora, J.P.; De Smet, C.; Brë
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A;Title: Structure, chromosomal localization, and expression of 12 genes of the MAGE fami
A;Reference number: 138659; MUID:95012457; PMID:7927540
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A;Experimental source: MAGE-5a antigen
A;Accession: 138664
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A; Residues: 1.124 CDEP2>
A; Cross-references: EMBL: U10690; NID: 9533520; PIDN: AAA68874.1; PID: 9533521
A; Cross-references: RAGE-5b antigen
A; Rxperimental source: MAGE-5b antigen
A; Note: these sequences seem to be incomplete with respect to other members of the superfix R; Traversari, C.; van der Bruggen, P.; Luescher, I.F.; Lurquin, C.; Chomez, P.; Van Pel, B; Exp. Med. 176, 1453-1457, 1992
A; Title: A nonapeptide encoded by human gene MAGE-1 is recognized on HLA-A1 by cytolytic A; Reference number: PH1294; MUID: 93018875; PMID: 1402688
                                                                                                                                                                                                                                                                                                                                                  A,Cross-references: UNIPROT:P25233; GB:M80840; NID:g200019; PIDN:AAA39805.1; PID:g200020
C;Superfamily: tumor associated protein MAGE
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              45 SSSFPSSFPSSSSSSSSSSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQ
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | : | : | : | : | : | | SDAVPVGIPPPASLAANLAGP--PCAPEGPMAAQQA-SPPPEERIEDVDPKI----LQQ
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      105 SDEGSSSOKEESPSTLQVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESV
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      165 IKNYEDHFPLLFSEASECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKT
                                                                                                                                                                                                                                                                                                                                                                                                                                         19.7%; Score 372; DB 2; Length 325;
29.6%; Pred. No. 4e-18;
tive 73; Mismatches 122; Indels
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A;Status: preliminary; translated from GB/EMBL/DDBJ
A;Molecule type: DNA
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A;Molecule type: DNA
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A;Residues: 125-133 <TRA1>
A;Experimental source: MAGE 5 protein
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A;Molecule type: mRNA
A;Residues: 1-325 <MAR>
JN0148
necdin, brain - mouse
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      66 LIPSTPREVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQVLPD 125
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A;Reference number: 217313
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A;Accession: Til678
A;Molecule type: DNA
A;Molecule type: DNA
A;Coss-references: UNIPROT:074346; EMBL:AL031536; NID:e1319499; PID:e1319505
A;Experimental source: strain 972h(-)
C;Genetics:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          hypothetical protein SPBC21D10.06c – fission yeast (Schizosaccharomyces pombe)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           6 KRORCMPEEDLOSOSETOGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSSSSCYP 65
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              6.7%; Score 126.5; DB 2; Length 948;
20.8%; Pred. No. 0.94;
tive 56; Mismatches 138; Indels 125; Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             25;
                                                                                                                                                                                                                                                                                                                                                                                                                                                       DB 2; Length 133;
                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match 11.8%; Score 222.5; DB 2; Length 1 Best Local Similarity 39.3%; Pred. No. 1.9e-08; Matches 57; Conservative 19; Mismatches 44; Indels
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A; Residues: 125-133 <TRA2>
A; Experimental source: MAGE 51 protein C; Genetics: A; Gene: GDB: MAGEA5; MAGES
A; Gross-references: GDB: 331120
A; Map position: Xq28-Xq28
A; Introns: #status absent
C; Superfamily: tumor associated protein MAGE
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Best Local Similarity 20.8%
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A, Accession: PH1300
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SEQUENCE FROM N.A.
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GenCore version 5.1.6
(c) 1993 - 2005 Compugen Ltd.
                                                                                                                                 Total number of hits satisfying chosen parameters:
                                                                                                                    1612378 seqs, 512079187 residues
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MAGB_HUMAN
QEZRZS
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Gapop 10.0 , Gapext 0.5
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MEDLINE=2388257; PubMed=12477932; DOI=10.1073/pnas.242603899;
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Strausberg R.L., Felingold E.A., Grouse L.H., Derge J.G.,
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Altschul S.F., Zeeborg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
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A stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
A promstein M.J., Uddin T.B., Toshiyuki S., Carninci P., Prange C.,
A Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
A Richards S., Worley D.M., Sodergren B.J., Lu X., Gibbs R.A.,
A Villalon D.K., Muzny D.M., Sodergren B.J., Lu X., Gibbs R.A.,
A Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
A Blakesley R.W., Touchman J.W., Green B.D., Dickson M.C.,
A Blakesley R.W., Touchman J.W., Green B.D., Dickson M.C.,
A blatchield Y.S.N., Krzywinski M.I., Skalaka U., Smailus D.E.,
A schnerch A., Schein J.B., Jones S.J.M., Marra M.A.,

"Generation and initial analysis of more than 15,000 full-length human FISSUE SPECIFICITY: Expressed in many tumors of several types, such as melanoma, head and neck squamous cell carcinoma, lung carcinoma and breast carcinoma, but not in normal tissues except for testes and placenta.
SIMILARITY: Contains 1 MAGE domain. (1) — SEQUENCE FROM N.A.

SEQUENCE FROM N.A.

MEDILINE-95012457; PubMed=7927540;

de Plaen E., Arden K., Traversari C., Gaforio J.J., Szikora J.-P.,

de Smet C., Brasseur F., van der Bruggen P., Lethe B., Lurquin C.,

"Structure, Chomez P., de Backer O., Cavenee W., Boon T.; genes of

the MAGE family."; Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi; Mammalia; Butheria; Primates; Catarrhini; Hominidae; Homo. Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
-I- FUNCTION: Not known, though may play a role in embryonal development and tumor transformation or aspects of tumor 01-NOV-1995 (Rel. 32, Created)
01-NOV-1995 (Rel. 32, Last sequence update)
05-UUJ-2004 (Rel. 44, Last annotation update)
Melanoma-associated antigen 10 (MAGE-10 antigen)
Name-MAGEAL0; Synonyms=MAGE10;
Homo sapiens (Human) 369 AA PRT; [mmunogenetics 40:360-369(1994) and mouse cDNA sequences."; STANDARD;

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MEDLINE=95012457; PubMed=7927540;
de Plaen B., Arden K., Travergari C., Gaforio J.J., Szikora J.-P.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          PAGGE HUMAN STANDARD; PRT; 319 AA.
943364;
01-304-1995 (Rel. 32, Created)
01-NOV-1995 (Rel. 32, Last sequence update)
05-JUL-2004 (Rel. 44, Last annotation update)
Melanoma-associated antigen 11 (MAGE-11 antigen).
Name-MAGEA11; Synonyms-MAGE11;
Name-MAGEA11; Synonyms-MAGE11;
Bukaryota; Metazoa; Chordae; Craniata; Vertebrata; Buteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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                                                                                                                                                                                                                                                                                                                                                                                        100.0%; Score 1891; DB 1; Length 369; 100.0%; Pred. No. 1e-115; tive 0; Mismatches 0; Indels 0
                                                                                                                                                                                                                                                                                                                                                          369 AA; 40766 MW; 16FA3301CAB716A6 CRC64;
                                                                                                                                                                                                                        H-Invus, ...
H-Invus, 300313; -..
InterPro; IPR002190; MAGE.
PFam; PF01454; MAGE; 1.
PROSITE; PS50838; MAGE; 1.
Antigen; Multigene family; Tumor antigen.
134 33 MAGE.
                                                                                                                                                                 EMBL; BC004105; AAH04105.1; -. PIR; I38659; 138659; 138659; Genew; HGNC:6797; MAGEALO.
H-INVDB; HIX0017116; -.
                                                                                                                                                   EMBL; U10685; AAA68869.1; -.
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ATGSFSYPE 369
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**XEXUE=SKINIS;**)
**REDLINE=2218825;** PubMed=12477932; DOI=10.1073/pnas.242603899;
**REDLINE=2218825;** PubMed=12477932; DOI=10.1073/pnas.242603899;
**Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G.,
**Alachul S.F., Zeeberg B., Buetcher, Schaefer C.F., Bhat N.K.,
**Alachul S.F., Zeeberg B., Buetcher, S.I., Wang J., Hasteh F.,
**Alachenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
**Rapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
**Rapleton M., Soares M.B., Peters G.J., Abramson R.D., Mullahy S.J.,
**Rab S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
**Rab S.S., McEwan P.J., McKernan K.J., Malek J.A., Gunarathe P.H.,
**Rab S.S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
**A Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
**A Richards S., Worley B., Sodergren B.J., Lu X., Gibbs R.A.,
**A Pathon D.K., Muzny D.M., Sodergren B.J., Lu X., Gibbs R.A.,
**A Pathon M. Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
**A Richards S. C., Gaimwood J., Schmutz J., Myers R.M.,
**Butterfield Y.S. N., Krzywinski M.I., Skalska U., Smailus D.E.,
**A Butterfield Y.S. N., Krzywinski M.I., Skalska U., Smailus D.E.,
**A Bard M. Madan A., Schein J.B., Jones S.J.M., Marra M.A.;
**A Generation and initial analysis of more than 15,000 full-length human
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@ib-sib.ch).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         61 SSCYPLIPSTPREVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
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TISSUE SPECIFICITY: Expressed in many tumors of several types, such as melanoma, head and neck squamous cell carcinoma, lung carcinoma and breast carcinoma, but not in normal tissues except
de Smet C., Brasseur F., van der Bruggen P., Lethe B., Lurquin C., Brasseur R., Chomez P., de Backer O., Cavenee W., Boon T.; "Structure, chromosomal localization, and expression of 12 genes of the MAGE family.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               22;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          and mouse cDNA sequences.";
Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
-!- PUNCTION: Not known, though may play a role in embryonal development and tumor transformation or aspects of tumor
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           81; Indels
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54.0%; Pred. No. 1.1e-51;
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DOMAIN 112 311 MAGE.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                for testes and placenta. SIMILARITY: Contains 1 MAGE domain.
                                                                                                                                                          Immunogenetics 40:360-369(1994)
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I38660; I38660.
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Pfam; PF01454; MAGE; 1.
PROSITE; PS50838; MAGE; 1.
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H-InvDB, HIX0022316; -.
MIM, 300344; -.
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Matches 183;
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       SOLUTION OF THE PROPERTY OF TH
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SEQUENCE FROM N.A.
TISSUE=Skin;
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                                                            Q9BUN9
                    RESULT 4
Q9BUN9
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                                        76 MPLEORSÓHCKPEEGLÓAÓEEDLGĽVGÁÓALQÁEEGEÁAFFSST------ 119
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Tashiro H., Yamazaki M., Watanabe K., Kumagai A., Itakura S., Fukuzumi Y., Fujimori Y., Komiyama M., Sugiyama T., Irie R., Otsuki T., Sato H., Wakamatsu A., Ishii S., Yamamoto J., Isono Y., Kawai-Hio Y., Saito K., Nishikawa T., Kimura K., Yamashita H., Matsuo K., Nakamura Y., Sekine M., Kikuchi H., Kanda K., Wagatsuma M., Murakawa K., Kanehori K., Takahashi-Fujii A., Oshima A., Sugiyama A., Kawakami B., Suzuki Y., Sugano S., Nagahari K., Masuho Y., Nagai K.,
                                                                                                                          TPREVIWEALNWMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYBFLWGP
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05-JUL-2004 (TrEMBLrel. 27, Last sequence update)
05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
Hypothetical protein FLJ45952.
Homo sapiens (Human).
Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                22;
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Submitted (JUL-2003) to the EMBL/GenBank/DDBJ databases.
EMBL; AK127849; BAC87161.1;
Interpro; IFR002190; MAGE.
Pfam; PF01454; MAGE; 1.
PROSITE; PS50838; MAGE; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     550838; MAGE; 1.
394 AA; 44276 MW; 36D7A81D8749D896 CRC64;
                                                                                                                                                                                                                                     RAHAETSKMKVLEYIANANGRDPTSYPSLYEDALREEGE 317
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ; Score 911; DB 2;
; Pred. No. 1.5e-51;
53; Mismatches 81;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     48.2%;
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                                                                                                                                                                                                                                                                                                                                                                PRELIMINARY;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         TISSUE=Placenta;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             NCBI_TaxID=9606;
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66
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Straubberg R.L., Feingold B.A., Grouse L.H., Derge J.G.,

Altauner R.D., Collins F.S., Magner L., Schaefer C.R., Schuler G.D.,

Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,

Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,

Antschenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,

Biatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,

Brownstein M., Soares M.B., Bonaldo M.F., Caarainci P., Prange C.,

Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,

Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,

Villalon D.K., Muzny D.M., Sodergren B.J., Lu X., Gibbs R.A.,

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Rachiquez A.C., Grimwood J., Schmutz J., Myers R.M., Butterfield Y.S.,

Krzywinski M.I., Skalska U., Smailus D.E., Schnerch A., Schein J.E.,

"Generation and initial analysis of more than 15,000 full-length human
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Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
NCBI_TaxID=9606;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             MEDLINE=22388257; PubMed=12477932; DOI=10.1073/pnas.242603899;
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56.5%; Pred. No. 1.3e-51;
ive 45; Mismatches 78; Indele
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Submitted (AUG-2001) to the EMBL/GenBank/DDBJ databases.
EMBL; BC002455; AAH02455.1; --
EMBL; BC007340; AAB36004.1; --
EMBL; BC012744; AAH12744.1; --
InterPro; IPR002190; MAGB.
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Submitted (FEB-2001) to the EMBL/GenBank/DDBJ databases
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           35214 MW; EA02C1FB42F6C080 CRC64;
                                                                Last sequence update)
Last annotation update)
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                                Created)
01-JUN-2001 (TrEMBLrel. 17, C. 01-JUN-2001 (TrEMBLrel. 17, L4 25-OCT-2004 (TrEMBLrel. 28, L6 Melanoma antigen, family A, 9.
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Best Local Similarity 56.5%;
Matches 188; Conservative
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                                                                                                                                                                                                                    Homo sapiens (Human)
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TISSUE-Duodenum;

X MEDLINE=22388257; PubMed=12477932; DOI=10.1073/pnas.242603899;

A Straubberg R.L., Feingold E.A., Grouse L.H., Derged J.G.,

A Klausner R.D., Collins P.S., Wagner L., Shemmen C.M., Schuler G.D.,

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B Rownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,

A Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,

B Bosak S.A., McKernan K.J., Malek J.A., Gunaratne P.H.,

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A Nilalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,

Fahey J., Helton E., Ketteman M., Madan A., Rodrigues S., Sanchez A.,

M Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,

B Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
SESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSBASECMLL 185
                    104 LESLFREALDEKVAELVRFLIRKYQIKEPVTKAEMLESVIKNYKNHFPDIFSKASECMQV 163
                                                                                            IWEALNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGPRAHAE 305
                                                                                                                                                                             MEDLINE-95012457; PubMed=7927540; de Plaen B., Arden K., Traversari C., Gaforio J.J., Szikora J.-P., de Plaen B., Brasseur F., van der Bruggen P., Lethe B., Lurquin C., Brasseur R., Chomez P., de Backer O., Cavenee W., Boon T., "Structure, chromesomal localization, and expression of 12 genes of the MAGE family.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  SEQUENCE FROM N.A.
MEDLINE=95369706; PubMed=7642112; DOI=10.1016/0378-1119(94)00680-Q; Imai Y., Shichijo S., Yamada A., Katayama T., Yano H., Itoh K.; "Sequence analysis of the MAGE gene family encoding human tumor-cejection antigens."; Gene 160:287-290(1995).
                                                                                                                                                                                                                                                                                                                                                                                                           01-NOV-1995 (Rel. 32, Created)
01-NOV-1995 (Rel. 32, Last sequence update)
25-OCT-2004 (Rel. 45, Last annotation update)
Melanoma-associated antigen 4 (MAGE-4 antigen) (MAGE-X2) (MAGE-41).
Name-MAGEA4; Synonyms-MAGE4;
Homb sapleans (Human).
Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   MEDLINE=94311935; PubMed=8037761;
Ding M., Beck R.J., Keller C.J., Fenton R.G.;
"Cloning and analysis of MAGE-1-related genes.";
Biochem. Biophys. Res. Commun. 202:549-555(1994)
                                                                                                                                                                                                                                    306 IRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEE 338
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P43358;
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This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the European Bioinformatics Institute. There are no restrictions on its mee by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               276
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               61 SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                98
Rodriguez A.C., Grimwood J., Schmutz J., Myers R.M.,
Butterfield Y.S.N., Krzywinski M.I., Skalaka U., Smailus D.E.,
Schnerch A., Schein J.E., Jones S.J.M., Marza M.A.,
"Generation and initial analysis of more than 15,000 full-length human
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         PROSITE; PS50838; MAGE; 1.
3D-structure; Antigen; Multigene family; Polymorphism; Tumor antigen.
DOMAIN 110 309 MAGE.
                                                                                                                                          progression.
TISSUE SPECIFICITY: Expressed in many tumors of several types, such as melanoma, head and neck squamous cell carcinoma, lung carcinoma and breast carcinoma, but not in normal tissues except for testes and placenta.
SIMILARITY: Contains 1 MAGE domain.
                                                                    and mouse cDNA sequences.";
Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
-!- FUNCTION: Not known, though may play a role in embryonal
                                                                                                                        development and tumor transformation or aspects of tumor
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    45.2%; Score 854; DB 1; Length 317; 51.9%; Pred. No. 6e-48;
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307 307 E -> Q (1n Ref. 2).
317 AA, 34929 MW, 3CPAC052B696257C CRC64;
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EMBL; U10688; AAA68872.1; -.
EMBL; U10340; AAA19007.1; -.
EMBL; D32077; BAA06843.1; -.
EMBL; BC017723; AAH17723.1; -.
PIR; 128661; 138661.
PDB; 114F; X-ray; C=230-239.
Genew; HGNC:6802; MAGEA4.
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Pfam; PF01454; MAGE; 1.
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H-InvDB; HIX0017105; -.
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Hopking R.F., Jordan H., Moore T., Max S.I., Wang J., Haidh F.,

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"Mannan A., Schain J.B., Marra M.A.,

Rochneration and initial analysis of more than 15,000 full-length human
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              progression.
TISSUB SPECIFICITY: Expressed in many tumors of several types, such as melanoma, head and neck squamous cell carcinoma, lung carcinoma and breast carcinoma, but not in normal tissues except for testes and placenta.
SIMILARITY: Contains 1 MAGE domain.
                                                                                                                                                                                                                                                                de Plaen E., Arden K., Traversari C., Gaforio J.J., Szikora J.-P., de Smet C., Brasseur F., van der Bruggen P., Lethe B., Lurquin C., Brasseur R., Chomez P., de Backer O., Cavenee W., Boon T.; "Structure, chromosomal localization, and expression of 12 genes of
                                                                                                                                                                  Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi,
Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo.
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MEDLINE=22388257; PubMed=12477932; DOI=10.1073/pnas.242603899;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              and mouse cDNA sequences.";
Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
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                                                                                                                                                                                                                                                                                                                                                                                                  Timms K.M., Bondeson M.L., Ansari-Lari M.A., Lagerstedt Nelson D.L., Pettersson U., Gibbs R.A., Submitted (SEP-1996) to the EMBL/GenBank/DDBJ databases.
                             MAG9 HUMAN STANDARD; PRT; 315 AA. P43362; 092910; 01-NOV-1995 (Rel. 32, Created) NO-1995 (Rel. 32, Last sequence update) 05-UTU-2004 (Rel. 44, Last annocation update) Melanoma-associated antigen 9 (MAGE-9 antigen) Homo sapiens (Human).
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or send an email to license@isb-sib.ch).
                                                                                                                                                                                                                                                     MEDLINE=95012457; PubMed=7927540;
                                                                                                                                                                                                                                                                                                                                    the MAGE family.";
Immunogenetics 40:360-369(1994).
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EMBL; BC0002351, AAH02351.1; -
PIR; 138668; 138668.
Genew; HGNC:6807; MAGEA9.
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                                                                                                                                                                                                                                  SEQUENCE FROM N.A.
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                 MAG9_HUMAN
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97 PAQLEFMFQBALKLKVAELVHFLLHYYRVKEPVTKAEMLESVIKNYKRYFPVIFGKASEF 156
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               157 MOVIFGTDVKEVDPAGHSYILVTALGLSCDSMLGDGHSMPKAALLIIVLGVILTKDNCAP 216
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         63 CYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQV 122
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                                                                                                                                                                                                                                                                                                                                         3 RAPKRORCMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSSS 62
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Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
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                                                                                                                                                                                                                                            DB 1; Length 315;
                                                                                                                                                                                                                                                                                                                                                                                      6 RSP---HCKPDEDLEAQGEDLGLMGAQEPTGEEERTTSSSD-----
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Zhu J., Feng Z., Guan X.;
Zhu J., Feng Z., Guan X.;
Submitted (MAY-2003) to the EMBL/GenBank/DDBJ databases.
EMBL; AY310325; AAP82171.1;
InterPro; IPR002190; MAGE.
Pfam; PP01454; MAGE; 1.
PROSITE; PS50838; MAGE; 1.
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Poly-Glu.
7FD2ED10D680D928 CRC64;
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Last annotation update)
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                                                                                                                                                                                                                                                                                    88;
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                                                                                                                                                                                                                                          Query Match
44.8%; Score 846.5; DB 1
Best Local Similarity 51.0%; Pred. No. 1.8e-47;
Matches 172; Conservative 48; Mismatches 88
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           tch
al Similarity 51.0%; Pred. No. 1.8e-47;
172; Conservative 48; Mismatches 88
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MIM; 300342; -.
InterPro; IPR002190; MAGB.
Pfam; PF01454; MAGB; 1.
PROSITE; PS50838; MAGE; 1.
Antigen; Multigene family; Tumor antigen.
DOMAIN 108 307 MAGE.
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01-OCT-2003 (TrEMBLrel. 25, Last sequen
01-MAR-2004 (TrEMBLrel. 26, Last annote
Melanoma antigen family A 9 (Fragment).
                                                                                                                                                                                               315 AA; 35088 MW;
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SEQUENCE
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rejection antigens."
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Matches 171; Conserv
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                                                            335 AMTS 338
                                                                                                                                                                                                                                                                         MAGE-4 protein.
                     ATGS
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Ganis familiaris (Dog).
Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi,
Mammalia, Butheria, Carnivora, Pissipedia, Canidae, Canis.
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SEQUENCE FROM N.A.

MEDLINE=Z0470599; PubMed=11019919;

MA Z., Khatlani T.S., Ohno K., Sasaki K., Inokuma H., Onishi T.;

"Clonding and sequencing of canine MAGE cDNA.";

Tissue Antigens 56:166-169(2000).

EMBL, AF187235; AAF01438.1;

InterPro; IPR002190; MAGE.

InterPro; IPR002190; MAGE.

PROSITE; PS50838; MAGE; 1.

SEQUENCE 378 AA; 42214 MW; AD09C6DC8E3C8D57 CRC64;
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49.2%; Pred. No. 3.8e-46;
ive 52; Mismatches 107; Indels
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Last sequence update)
Last annotation update)
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(TrEMBLrel. 13, I
(TrEMBLrel. 23, I
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01-MAR-2003
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Best Local 9
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126 SESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEASECMLL 185
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     186 VFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILLILSIIFIEGYCTPEEV 245
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            246 IWEALNMWGLYDGMEHLIYGEPRKILIQDWVQENYLEYRQVPGSDPARYEFLWGPRAHAE 305
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       24; Gaps
                                                                                                                                                                                                                                                                                                                                                      MEDINE-95369706; PubMed=7642112; DOI=10.1016/0378-1119(94)00680-Q; Imai Y., Shichijo S., Yamada A., Katayama T., Yano H., Itoh K.; "Sequence analysis of the MAGE gene family encoding human tumor-
                                                                                                                                                                            Homo sapiens (Human).

Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Butheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 317;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            6 KSOHCKPEEGVEAQEBALGLVGAQAPTTEEQEAAVSSSS-----
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    86; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             317 AA; 35044 MW; 9B9477253FE307C4 CRC64;
                                                                                Last sequence update)
Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Last sequence update)
Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    306 IRKMSLLKFLAKVNGSDPRSFPLWYBEALKDEEE 339
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ::|:::|| ::|| 282 TSYVKVLEHVVRVNARVRIAYPSLREAALLEEEE 315
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 43.7%; Score 827; DB 2; 51.2%; Pred. No. 3.5e-46;
317 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       53; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Ź
                                                      Created)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             01-MAR-2003 (TrEMBLrel. 23, C3
01-MAR-2003 (TrEMBLrel. 23, L6
01-UDN-2003 (TrEMBLrel. 24, L6
Melanoma antigen, family B, 4
Name=MAGEB4;
Homo sapiens (Human)
                                                01-NOV-1996 (TrEMBLrel. 01, 01-NOV-1996 (TrEMBLrel. 01, 01-MAR-2003 (TrEMBLrel. 23,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Gene 160:287-290(1995).
EMBL, D32075; BAA06841.1; -InterPro., IPR002190; MAGE.
Pfam; PF0454; MAGE; I.
PROSITE; PS50838; MAGE; I.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Conservative
PRELIMINARY;
                                                                                                                                                                Name=melanoma antigen-4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               PRELIMINARY;
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ON SERVICE SER

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EMBL; U93163; AAC23619.1; -. EMBL; AC005185; AAD10637.1; Genew; HGNC:6811; MAGEB4.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           PROSITE; PS50838; MAGE; 1.
Antigen; Multigene family.
DOMAIN 109
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                InterPro, IPR002190; MAGE.
Pfam; PF01454; MAGE; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Local Similarity
les 177; Conservat
                                                                                                                            Homo sapiens (Human)
                                                                                                                                                                                                                                                                                                                                                                                                                                                               SEQUENCE FROM N.A.
15-DEC-1998 (Rel.
15-DEC-1998 (Rel.
05-JUL-2004 (Rel.
                                                                                                                                                                                                    NCBI_TaxID=9606;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           MIM; 300153;
                                                                                                       Name=MAGEB4;
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     셤
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                                                                                          Krausberg R.L., Feingold E.A., Grouse L.H., Derge J.G., Gruler G.D., Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G., Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G., Schuler G.D., Altschul S.F., Zeeberg B., Buerow K.H., Schaefer C.F., Bhat N.K., Altschul S.F., Zeeberg B., Buerow K.H., Schaefer C.F., Bhat N.K., Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Hsieh F., Dittchenko L., Marusiaa K., Farmer A.A., Rubin G.M., Hong L., Stapleton M., Soares M.B., Bonaldo M.F., Carnhorl P., Frange C., Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J., Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H., Richards S., Worley K.C., Hale S., Garria A.M., Gay L.J., Hulyk S.W., Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A., Halton E., Ketreman M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G., Mhiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G., Multing M., Madan A., Young A.C., Schmutz J., Myers R.M., Butterfield Y.S., Norley R.M., Touchman J.W., Green E.D., Dickson M.C., Grimwood J., Schmutz J., Myers R.M., Butterfield Y.S., Jones S.J., Marra M.A., Smailus D.E., Schmerch A., Schein J.E., Schmert A., Schein J.E., Schmer Human D.M., Schmutz J., Myers R.M., Butterfield Y.S., Warra M.A., Schein J. Schmutz J., Myers R.M., Butterfield Y.S., Warra M.A., Schein B.D., Schmutz J., Myers R.M., Butterfield Y.S., Warra M.A., Schein B.D., Schmutz J., Myers R.M., Butterfield Y.S., Warra M.A., Schmitting M. Marra M.A., Smailus D.E., Schmerch A., Schein J. B., Schmer Human D.M., Schmut B., Schmet L., Schmut B., Schmet Human B., Warra M.A., Schmitting M. Marra M.A., Smailus D.E., Schmet Human D.M., Schmitting M. 
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      61 SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  96 QASTSTERSLKDSLTRKTKOMLVQPLLYKYKOMKEPTIKAEMLKIISKKYKEHPPEIFRKVS 155
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILISIIFIEGYC 240
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    215 AREEEIWEFLMMLGIYDGKRHLIFGEPRKLIIQDLVQEKYLEYQQVPNSDPPRYQFLWGF 274
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKBPITKABILESVIKNYBDHFPLLFSEAS 180
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            241 TPEEVIWEALNWMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP 300
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 RAHAEIRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEEER--AQDRIATTDDTTAMASAS 358
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           1 MPRGQKSKLRAREKRQRTRGQTQDLKVGQ-PTAAEKEESPS------PSSSVLRDTASS 52
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   53 SLAFG-IPQEPQR-----EPPTTSAAAAMSCTG------SDKGDESQDEENASSS 95
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 1 MPRAPKRQRCMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSSFSSSSSS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          43.0%; Score 813; DB 2; Length 346; 47.6%; Pred. No. 3.3e-45; Live 60; Mismatches 105; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Strausberg R.; Submitted (JUN-2002) to the EMBL/GenBank/DDBJ databases. Submitted (JUN-2002) to the EMBL/GenBank/DDBJ databases. INTERPROS 1907 MAGE.

InterPro; IPR04254; MAGE; 1.

PR081TR; PS05038; MAGE; 1.

SEQUENCE 346 AA; 38981 MW; 724359116DBC8BE9 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903 (2002).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best_Local Similarity 47.6
Matches 175; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              and mouse cDNA sequences.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      SEQUENCE FROM N.A.
       NCBI_TaxID=9606;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                rissum=Testis;
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MGB4 HUMAN
ID MGB4 HUMAN
AC 015481;
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346 AA

STANDARD;

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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            116 SPSTLQVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKRPITKABILESVIKNYEDHFPLL 175
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   176 PSEASECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIF 235
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Muzny D., Arensonn A.D., Adams C., Brundage E., Bunac C., Carvelli K., Chacko J., Chen J., Di W., Ding Y., Dugan S., Durbin J., Forcum J., Ganesh R., Garcia C., Goodman M., Gorrell J.H., Haywood M., Hernandez J., Jackeon L., Jin S., Kampal R., Karpathy S., Kovar C., Leal B., Li Y., Lichtearge O., Liu W., Logan O., Lu J., Ly T., Martinez C., Oswal G., Perez L., Rashid N.D., Rowland K., Savage L., Scherer S.E., Shen H., Simon M., Stovall K., Timms K.M., Todd J., Vo Q., Williamson A., Worley K.C., Yu W., Chinault C., Nelson D., Gibbs R.A.;
Submitted (OCT-1998) to the EMBL/GenBank/DDBJ databases.
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                                                                                                                                                                                                                                                                                                                                                                         SEQUENCE PROM N.A.

BEDLINE-98110575. PubMed=9441743; DOI=10.1006/geno.1997.5052;

Lurquin C., de Smet C., Brasseur F., Muscatelli F., Martelange V., de Plaen E., Brasseur R., Monaco A.P., Boon T.;

TWO members of the human MAGEB gene family located in Xp21.3 are "Typessed in tumors of various histological origins.";

Genomics 46:397-408(1997).
                                                                                                                                                                                             Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi,
Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo.
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15-DEC-1998 (Rel. 37, Last sequence update)
05-JUL-2004 (Rel. 44, Last annotation update)
Melanoma-associated antigen B4 (MAGE-B4 antigen)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   42.9%; Score 812; DB 1; 47.5%; Pred. No. 3.8e-45;
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Eutheria; Primates; Catarrhini; Hominidae; Homo.
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                               SEQUENCE FROM N.A.
                                                                                                                   SEQUENCE FROM N.A.
           NCBI_TaxID=9606;
                                                                                     human melanoma
                                                                                                                                                                                                                                                                                                                                                                                          MUTAGENESIS
                                                                                                                                                                                                                                                                                                                                                                     Submitted
                                                                                                                                                                                                                                                                                            and man.
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                                 210 FLWGPRAHAETSKWKVLEFLAKVNDTTPNNFPLLYEBALRDEEERAGARPRVAARRGTTA 329
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             304
 210 LNGNCAREEEIWEFLNMLGIYDGKRHLIFGEPRKLITQDLVQEKYLEYQQVPNSDPPRYQ 269
                                                                                                                                                                                                                                                                                                                                                                                                                                              124
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKABILESVIKNYEDHFPLLFSEASECML 184
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                220 EIWEELSVWEVYDGREHSAYGEPRKLLTQDLVQEKYLEYRQVPDSDPARYEFLWGPRALA 279
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                   66
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   245 VIWEALNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGPRAHA
                      FLWGPRAHAEIRKMSLLKFLAKVNGSDPRSFPLWYBEALKDEBER--AQDRIATTDDTTA
                                                                                                                                                                                                                                                                                                                                                                                                    6 KRORCMPEEDLOSOSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSSSCYP
                                                                                                                                                                                                                                                                                                                                                                                                                        6 RSÓHCKPÉBGLEARGEALGLVGAQAPATEEQEAASSST-------
                                                                                                                                                                                                                                                                                                                                                                                                                                              LIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPL-DQSDEGSSSQKEESPSTLQVLP
                                                                                                                                                                                                                                                                                                                                                                                                                                                          LVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGYCTPEE
                                                                                                                                                                                                                                                                                                                                                                                Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             MAG1 HUMAN STANDARD; PRT; 309 AA.
P4335; 000346;
01-NOV-1995 (Rel. 32, Created)
01-NOV-1995 (Rel. 32, Last sequence update)
05-JUL-2004 (Rel. 44, Last annotation update)
Melanoma-associated antigen 1 (MAGE-1 antigen) (Antigen MZ2-E).
Name=MAGEAl; Synonyms=MAGEI, MAGEIA;
HOmo aapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
                                                                                                                                                                                                       Homo sapiens (Human).
Vertebrata; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Butheria; Primates; Catarrhini; Hominidae; Homo.
VCBI_TaxID=9606;
                                                                                                                                                                                                                                                                                                                                                                               27;
                                                                                                                                                                                                                                                                                                                                                           Length 316;
                                                                                                                                                                                                                                                                                                                                                                                83; Indels
                                                                                                                                                                                                                                                                    TISSUE-Hepatocellular carcinoma;
Sui Y., Ye J., Nu W.;
Submitted (NOV-201) to the EMBL/GenBank/DDBJ databases.
EMBL; AF443295; AAL37897.1; --
Pfam; PF01454; MAGE; 1.
PROSITE; PS50838; MAGE; 1.
                                                                                                                                                                                                                                                                                                                           350838; MAGE; 1.
316 AA; 35409 MW; A463A9A740A089DF CRC64;
                                                                                                                                                             Created)
Last sequence update)
Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     EIRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
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ETSYVKVLEYVIKVSARVRFFFPSLREALREEEE 314
                                                                                                                                                                                                                                                                                                                                                        Score 811.5; DB 2
Pred. No. 3.6e-45;
                                                                                                                                          316 AA
                                                                                                                                                                                                                                                                                                                                                                                49; Mismatches
                                                                                                                                          PRT;
                                                                                                                                                                                                                                                                                                                                                           42.9%;
52.5%;
                                                                                                                                                             01-MAR-2002 (TrEMBLrel. 20, 01-MAR-2002 (TrEMBLrel. 20, 01-MAR-2003 (TrEMBLrel. 23,
                                                              354 MASASSSATGSFS 366
                                                                                    MTSAYSRATSSSS 342
                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity 52.5
Matches 176; Conservative
                                                                                                                                          PRELIMINARY;
                                                                                                                                                                                             Tumor antigen MAGE-N.
                                                                                                                                                                                                                                                           SEQUENCE FROM N.A.
                      962
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                                                                                                                                                                                                                                                                                                                                      SEQUENCE
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MAG1 HUMAN
ID MAG1
AC P4335
DT 01-NO
DT 05-JU
DE Melan
GN Name=
GN Name=
OC Bukar
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This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (see http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              MEDLINE=20314869; PubMed=10854409; DOI=10.1101/gr.10.6.758;
MEDLINE=20314869; PubMed=10854409; DOI=10.1101/gr.10.6.758;
Mallon A.M., Platzer M., Bates R., Gloeckner G., Botcherby M.,
Nordsiek G., Strivens M.A., Kloschis P., Dangel A., Cunningham D.,
Straw R., Weston P., Hunter C., Gilbert M., Fernando S., Goodall K.,
Kerry G., Greystrong J.S., Clark D., Goerdes M., Blechschmidt K.,
Rump A., Hinzmann B., Mundy C.R., Miller W., Poustka A., Herman G.E.,
Rhodes M., Denny P., Rosenthal A., Brown S.D.M.;
"Comparative genome sequence analysis of the Bpa/Str region in mouse
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    MEDIINE-55112905; PubMed-7927954; Schultz-Thater E., Juretic A., Dellabona P., Luscher U., Siegrist W., Harder F., Heberer M., Zuber M., Spagnoli G.C.; MAGE-1 gene product is a cytoplasmic protein."; Int. J. Cancer 59:435-419(1994).
MEDLINE=92086861, PubMed=1840703; van der Bruguin C., de Plaen E. van der Bruggen P., Traversan C., Chomez P., Lurquin C., de Plaen E. van den Eynde B., Knuth A., Boon T.; Son T.; "A gene encoding an antigen recognized by cytolytic T lymphocytes on
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        FUNCTION: Not known, though may play a role in embryonal development and tumor transformation or aspects of tumor progression. Antigan recognized on a melanoma by autologous cytolytic T lymphocytes subcellular LOCATION: Cytoplasmic.

TISSUE SPECIFICITY: Expressed in many tumors of several types, such as melanoma, head and neck squamous cell carcinoma, lung carcinoma and breast carcinoma, but not in normal tissues except for testes. Never expressed in kidney tumors, leukemias and
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              TISSUE=Blood;
MEDLINE=44157413; PubMed=8113684;
Gaugler B., van den Bynde B., van der Bruggen P., Romero P.,
Gaforio J.J., de Plaen E., Lethe B., Brasseur F., Boon T.;
"Human gene WAGE-3 codes for an antigen recognized on a melanoma by
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Υ.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       polymorphism of MAGE-1 gene in Chinese people."; ttted (SEP-2002) to the EMBL/GenBank/DDBJ databases.
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Chen H., Wang L., Mei M., Qin L., Cong X., Xu J., Wei
                                                                                                                                                                                                                                                                                                                                                                                                                          MEDLINE=94311935; PubMed=8037761;
Ding M., Beck R.J., Keller C.J., Fenton R.G.;
"Cloning and analysis of WdE-1-related genes.";
Blochem. Biophys. Res. Commun. 202:549-555(1994)
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J. Exp. Med. 179:921-930(1994).
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                                                                                                                                                                                                                                        Science 254:1643-1647(1991).
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121 QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS 180
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      241 TPEEVIWEALNWMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP 300
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Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H., Richards S., Worley K.C., Hale S., García A.M., Gay L.J., Hulyk S.W., Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A., Hulyk S.W., Muzly D.M., Ketteman M., Madan A., Rodrigues S., Sanchez A., Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G., Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C., Surkertleid Y.S., Krzywinski M.I., Skalska U., Schmutz J., Myers R.M., Butterfield Y.S., Krzywinski M.I., Skalska U., Smailus D.E., Schnerch A., Schein J.E., Generation and initial analysis of more than 15,000 full-length human and mouse cDMA sequences.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TISSUE=Breast;
MEDLINE=22388257; PubMed=12477932; DOI=10.1073/pnss.242603899;
Strausberg R.L., Felngold E.A., Grouse L.H., Derge J.G.,
Klauener R.D., Collins E.S., Wagner L., Shemmen C.M., Schuler G.D.,
Altschul S.P., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Heleh F.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi,
Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo.
NCBI_TaxID=9606;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Strausberg R.; Strausberg R.; Strausberg R.; Submitted (AUG-2001) to the EMBL/GenBank/DDBJ databases Submitted (AUG-2001) to the EMBL, BC013098; AAH13098.1; -. InterPro; IPR002190; MAGE.; I. Pfam; PF01454; MAGE; 1. PR051TE; PS160818; MAGE; 1. SEQUENCE 314 AA; 35024 MW; A94F16247D1BFBCO CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            301 RAHABIRKMSLLKFLAKVNGSDPRSFPLWYBBALKDEBE 339
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Last annotation update)
                                                                                                                                                                                                 Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002)
                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
42.6%; Score 806.5; DB 2;
Best Local Similarity 49.6%; Pred. No. 7.6e-45;
Matches 168; Conservative 54; Mismatches 92;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       PRELIMINARY;
                                                                                                                                                                                                                                   SEQUENCE FROM N.A.
TISSUE=Adrenal cortex;
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Q6P448;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                98 FRAVITKKVADLVGFLLLKYRAREPUTKAEMLESVIKNYKHCFPEIFGKASESLQLVFGI 157
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      70 TPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQVLPDSESL 129
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MEDLINE-22388257; PubMed=12477932; DOI=10.1073/pnas.242603899;
Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G.,
Klauener R.D., Collins F.S., Wagner L., Schemen C.M., Schuler G.D.,
Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Hsieh F.,
Diatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz
Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Frange C.,
Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
                                                                                                                                                                                                                                                                                                                                                                                 Gaps
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Mammalia; Butheria; Primates; Catarrhini; Hominidae; Homo.
NCBI_TaxID=9606;
                                                                                                                                                                                                                                                                                                                                                                                 32;
                                                                                                                                                                                                  T -> A (probable polymorphism).
/FIId-VAR 004283.
R -> Q (in dbSNP:2008144).
/FIId-VAR 011737.
D-AR Abolishes HLA-Al binding.
Y-AR Abolishes HLA-Al binding.
                                                                                                                                                                                                                                                                                                                                          Query Match 42.7%; Score 807; DB 1; Length 309; Best Local Similarity 53.6%; Pred. No. 6.9e-45; Matches 177; Conservative 40; Mismatches 81; Indels
                                                                                                                                      Antigen; Multigene family; Polymorphism; Tumor antigen. DOMAIN 102 301 MAGR.
                                                                                                                                                                                                                                                                                                         544EEB1F9F4E9D33 CRC64;
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Last sequence update)
Last annotation update)
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169 169 Y
309 AA; 34342 MW;
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       EMBL; AY148486; AAN62752.1;
                                                                                        InterPro; IPR002190; MAGE.
Pfam; PF01454; MAGE; 1.
PROSITE; PS50838; MAGE; 1.
                           MAGEA1
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                    Genew; HGNC:6796; MAGEAL
H-InvDB; HIXO017126; -.
MIM; 300016; -.
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MUTAGEN
SEQUENCE
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Indele

Length 314;

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Distribution L., Marusina K., Farmer A.A., Rubin G.M., Hong L., Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.B., Brownstein M.J. Usdin T.B., Toshiyuki S., Carninci P., Prange C., A Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C., A Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J., McKernan K.J., Malek J.A., Gunaratne P.H., Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W., A Hilahon D.K., Muzuy D.W., Sodergren B.J., Lu X., Gibbs R.A., Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G., Mitching M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G., Rodriguez A.C., Grimwood J., Schmutz J., Myers R.W., Buterfield Y.S., Krzywinski M.I., Skalska U., Smailus D.E., Schnerch A., Schein J.E., Jones S.J., Marra M.A.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ë,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       156 EYLQLVPGIEVVEVVPISHLYILVTCLGLSYDGLLGDNQVMPKTGLLIIVLAIIAIEGDC 215
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   216 APEEKIWEEJ,SMLEVFEGREDSVFAHPRKLIAMQDLVQENYLEYRQVPGSDFACYEFLWGP 275
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         61 SSCYPLIPSTPREVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    121 QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS 180
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGYC 240
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         241 TPEEVIWEALNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP 300
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              1 MPRAPKRQRCMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSS 60
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
42.5%; Score 804.5; DB 2; Length 3
Best Local Similarity 49.6%; Pred. No. 1e-44;
Matches 168; Conservative 53; Mismatches 93; Indels
                                                                                                                                                                                                                                                                                                                                                                         Strausberg R.; Strausberg R.; Submitted (DEC-2003) to the EMBL/GenBank/DDBJ databases. Submitted (DEC-2003) to the EMBL/GenBank/DDBJ databases. InterPro; IPR002190; MAGE.

Pfam; PF01454; MAGE; 1.

PROSITE; PSSO038; MAGE; 1.

SEQUENCE 314 AA; 35026 MW; 28810BE358FA3FE6 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              301 RAHABIRKMSLLKFLAKVNGSDPRSFPLWYBEALKDEBE 339
                                                                                                                                                                                                                                                                          and mouse cDNA sequences.";
Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002)
                                                                                                                                                                                                                                                                                                                                       SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                         TISSUE=Breast;
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completed: November 15, 2005, 15:43:46 ne : 218.738 secs Search con Job time :

3, Appli 8202, Ap 26, Appl 5, Appli

Sequence 26, Appl Sequence 11171, A Sequence 11172, A Sequence 3, Appl Sequence 26, Appl Sequence 26, Appl Sequence 5, Appl Sequence 6, Appl Sequence 6, Appl Sequence 6, Appl Sequence 11308, Appl

Appl Appli

Sequence (Sequence (Sequence (

Scoring table:

Searched:

Minimum DB E Maximum DB E

Database

Perfect score:

Sequence:

OM protein

Run on:

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Length 369;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Sequence 4, Application US/08773870;
Patent No. 5912143;
GENERAL INFORMATION:
APPLICANT: Bandman, Olga
APPLICANT: Goli, Surya K.
TITLE OF INVENTION: NOVEL HUMAN MAGE-LIKE PROTEIN
NUMBER OF SEQUENCES:
CORRESPONDENCE ADDRESS:
ADDRESSEE: Incyte Pharmaceuticals, Inc.
STREET: 3174 Porter Drive
CITY: Palo Alto
STATE: CA
COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match 100.0%; Score 1891; DB 2; Best Local Similarity 100.0%; Pred. No. 1.2e-172; Matches 369; Conservative 0; Mismatches 0;
                 US-09-392-714-26
US-09-270-437D-2
US-09-949-016-11171
US-09-949-016-11172
US-09-748-016-11172
US-09-949-016-8202
US-09-948-4310-26
US-09-468-4330-24
US-09-362-123A-6
US-09-362-123A-6
US-09-949-016-6944
US-09-949-016-6944
US-09-949-016-6944
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 COMPUTER READBLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: Diskette
COMPUTER: Diskette
OFFRATING SYSTEM: DOS
SOFTWARE: FRANCHING DOS
SOFTWARE: FRANCHING DATA:
APPLICATION NUMBER: US/08/773,870
FILING DATE: Herewith
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US/08/773,870
FILING DATE: HEREWITH
APPLICATION NUMBER: 36,749
REGISTRATION NUMBER: 36,749
REFERENCE/DOCKET NUMBER: 36,749
                                                                                                                                                                                                                                                                                                                                                          JS-09-621-976-6400
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US-09-662-831-1
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SEQUENCE CHARACTERISTICS:
LENGTH: 369 amino acids
TYPE: amino acid
STRANDEDNESS: single
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TOPOLOGY: 1110C.
IMMEDIATE SOURCE:
LIBRARY: Genbank
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8001264566666444444
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8559, Ap
22, Appli
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8188, Ap
                                                                                                                                                                    (without alignments)
462.579 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                                                                                ; Search time 59.5476 Seconds
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Sequence 19,
Sequence 19,
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Sequence 1
Sequence 1
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Sequence 8
Sequence 8
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Sequence
Sequence
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Sequence
Sequence
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1 MPRAPKRQRCMPEEDLQSQS......DTTAMASASSSATGSFSYPE
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    /cgn2_6/ptodata/1/iaa/5A_COMB.pep:*
/cgn2_6/ptodata/1/iaa/5B_COMB.pep:*
/cgn2_6/ptodata/1/iaa/6A_COMB.pep:*
/cgn2_6/ptodata/1/iaa/ReCTUS_COMB.pep:*
/cgn2_6/ptodata/1/iaa/PCTUS_COMB.pep:*
                     GenCore version 5.1.6
(c) 1993 - 2005 Compugen Ltd.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               US-09-949-016-8188
US-08-455-167A-24
US-08-845-167A-24
US-08-845-1618-10
US-08-627-820-24
US-09-96-281B-10
US-09-949-016-6574
US-09-949-016-6574
US-09-949-016-6574
US-09-949-016-6574
US-09-949-016-6574
US-09-949-016-6574
US-09-948-015-2
US-09-96-215-2
US-09-96-215-2
US-09-96-215-2
US-09-96-2181-19
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                                                                                                                                                                                                                                                                                                                                                                                   513545 segs, 74649064 residues
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                                                                                                                                                November 15, 2005, 15:33:16
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                                                                         - protein search, using sw model
                                                                                                                                                                                                                                                                                                                                         Gapop 10.0 , Gapext 0.5
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                    seq length: 0
seq length: 200000000
                                                                                                                                                                                                                                    US-09-856-812B-1
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Match Length DB
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1142
1142
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911 807 807 807 807 807 807 807 807 793.5 793.5 783.5 783.5 783.5

Score

So.

Result

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Gaps

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241 TPEEVIWEALNAMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP 300
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159 VCMQLLFGIDVKEVDPTSHSYVLVTSLNLSYDGIQCNEQSMPKSGLLIIVLGVIFMEGNC 218
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               CKPEEALEAQQEALGLVCVQA------ATSSSS-------PLVLG 41
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 309;
                                                                                                                                                                                                                                                              Sequence 24, Application US/08465167A

Patent No. 5750395

GENERAL INFORMATION:
APPLICANT: Fives, John D.
APPLICANT: Sette, Alessandro D.
APPLICANT: Sette, Alessandro D.
TITLE OF INVENTION: INMUNOGENIC PEPTIDES (as amended)
NUMBER OF SEQUENCES:
CORRESPONDENCE ADDRESS:
ADDRESSEE: Townsend and Crew LLP
                                                                                                                        301 RAHABIRKMSLLKFLAKVNGSDPRSFPLWYBBALKDBEE 339
                                                                                                                                              RAHAETSKWKVLEYIANANGRDPTSYPSLYEDALREEGE 317
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         MEDIUM TYPE: Floppy disk
COMPUTER: IEM PC Compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.25
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      42.7%; Score 807; DB 1;
ilarity 53.6%; Pred. No. 7.6e-69;
Conservative 40; Mismatches 81
                                                                                                                                                                                                                                                                                                                                                                                                                                                                            STREET: Two Embarcadero Center, 8th Floor
CITY: San Francisco
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/465,167A
FILING DATE: 05-JUN-1995
CLASSIFICATION: 435
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                FILING DATE: 05-UUN-1995
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/103,623
FILING DATE: 06-AUG-1993
ATTORNEY/AGENT INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      NAME: Parmelee, Steven W.
REGISTRATION NUMBER: 31,990
REPERENCE/DOCKET NUMBER: 14137
TELECOMMUNICATION INPORMATION:
TELEPAN: 206-467-9600
TELEPAX: 415-576-0300
INPORMATION FOR SEQ ID NO: 24:
SEQUENCE CHARACTERISTICS:
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Matches 177; Conserv
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Sequence 8188, Application US/09949016

Patent No. 6812339

GENERAL INFORMATION:
APPLICANT: VENTER, J. Craig et al.
TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED
TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED
TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF
PILE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF
CURRENT APPLICATION NUMBER: US/09/949,016

PRIOR APPLICATION NUMBER: 60/241,755
PRIOR PILING DATE: 2000-10-20

PRIOR PILING DATE: 2000-10-03

PRIOR PILING DATE: 2000-10-03

PRIOR FILING DATE: 2000-10-03

NUMBER: OF SEQ ID NOS: 207012

SEQ ID NOS: 207012

SEQ ID NO 9188
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                                                                                                                                                                                                                                                                                            ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILLLILSIIFIEGYC 240
                                                                                                                                                                                                                                                                                                                                                                                                                                 RAHAEIRKWSLLKFLAKVNGSDPRSFPLWYEEALKDEEERAQDRIATTDDTTAMASASSS 360
                                                                                                                                                                                                                                                                                                                                                                                                                                                       SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
                                                                                                                                                                                QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS 180
                                                                                                                                                                                                                                                                  ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGYC 240
                                                                                                                                                                                                                                                                                                                                                  TPEEVIWEALNWMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP 300
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  99 PDLIDPESFSQDILHDKIIDLVHLLIRKYRVKGLITKAEMLGSVIKNYEDYFPEIFREAS 158
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             181 ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILISIIFIEGYC 240
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             61 SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
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                                             MPRAPKRQKROMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSS
                                                                                                                                                                                                          QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKŒPITKAEILESVIKNYEDHFPLLFSEAS
                                                                                                                                                                                                                                                                                                                                                                           241 TPEEVIWEALNAMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP
                                                                                                   SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL
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48.2%; Score 911; DB 4; Length 31
Best Local Similarity 54.0%; Pred. No. 8.6e-79;
Matches 183; Conservative 53; Mismatches 81; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ATGSFSYPE 369
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ORGANISM: Human
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US-09-949-016-8188
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US-09-949-016-8188
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130 PRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEASECMLLVFGI 189
98 FRAVITKKVADLVGFLLLKYRAREPVTKAEMLESVIKNYKHCFPEIFGKASESLQLVFGI 157
                                                                                      DVKEADPTGHSYVLVTCLGLSYDGLLGDNQIMPKTGPLIIVLVMIAMEGGHAPEEEIWEE 217
                                                                                                                                                 250 LINMMGLYDGMEHLIYGEPRKILITQDWVQENYLEYRQVPGSDPARYEFLWGPRAHAEIRKM 309
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                                                                                                                                                                                                                                                                                                                                                                      Sequence 10, Application US/08845528C

Sequence 10, Application US/08845528C

Batent No. 6027924

GENERAL INFORMATION

APPLICANT: LUCAS, Sophie;

APPLICANT: BOON-FALLER, Thierry

TITLE OF INVENTION: ISDIATED NUCLEIC ACID MOLECULE CODING FOR TUMOR ITLE OF INVENTION: THEREOF

TITLE OF INVENTION: THEREOF

TITLE OF INVENTION: THEREOF

NUMBER OF SEQUENCES: 14

CORRESPONDENCE ADDRESS:

ADDRESSEE: Felfe & Lynch

STREET: 805 Third Avenue

CITY: New York City

STATE: New York

COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10 CMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSSSSCYPLIPS 69
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 TLEEVPTAGST-DPPQSPQGASAFPTTINFTRQRQPSEGSSSREEEGPSTSCIL---ESL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          32; Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          81; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Diskette, 3.5 inch, 360 kb storage
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  42.7%; Score 807; DB 3; 53.6%; Pred. No. 7.6e-69;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            40; Mismatches
                                                                                                                                                                                                                                   310 SLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
                                                                                                                                                                                                                                                            SOFTWARE: Wordperfect
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/845,528C
FILING DATE: April 25, 1997
CLASSIFICATION: 4335
ATTORNEY/AGENT INFORMATION:
NAME: MARY ATHO SCHOÓLGÍGI
REGISTRATION NUMBER: 36,669
REFERENCE/DOCKET NUMBER: LUD 5455
TELECOMMUNICATION: INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        , TOPOLOGY: linear 'S-08-845-528C-10
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TELEPHONE: (212) 688-9200
TELEPAX: (212) 888-984
INPORMATION FOR SEO ID NO: 10
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Matches 177; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   COMPUTER READABLE FORM:
MEDIUM TYPE: Diskett
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             COMPUTER: IBM PS/2
OPERATING SYSTEM:
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Best Local Similarity
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                                                                                                            158
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                                                  130 PRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEASECMLLVFGI 189
                          DVKKVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILLILSIIFIEGYCTPBEVIWEA 249
                                                                                                            250 LNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGPRAHAEIRKM 309
                                                                                                                                                 218 LSVMEVYDGREHSAYGEPRKLLTQDLVQEKYLEYRQVPDSDPARYEFLWGPRALAETSYV 277
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10 CMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSSSSSCYPLIPS 69
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Sequence 10, Application US/08993118
Patent No. 5997872
GENERAL INFORMATION:
APPLICANT: LUCAS, Sophie;
APPLICANT: BOON-FALLEUR, Thierry
TITLE OF INVENTION: ISOLATED NUCLEIC ACID MOLECULE CODING FOR TUMOR TITLE OF INVENTION: RESECTION ANTIGEN PRECURSOR MAGE-C1 AND USES TITLE OF INVENTION: THEREOF
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    32;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       COUNTAL 10022
ZIP 10022
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette, 3.5 inch, 360 kb storage
"MEDIUM TYPE: Diskette, 3.5 inch, 360 kb storage
"MEDIUM TYPE: Diskette, 3.5 inch, 360 kb storage
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278 KVLEYVIKVSARVRFFFPSLREAALREEEE 307
                                                                                                                                                                                              310 SLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
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PRIOR APPLICATION DATA:
APPLICATION NUMBER: US/08/845,528
FILING DATE: April 25, 1997
ATTORNEY/AGENT INFORMATION:
NAME: Mary Anne Schoffeld
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             REGISTRATION NUMBER: 36,669
REFERENCE/DOCKET NUMBER: LUD 5455
TELECOMMUNICATION INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       US/08/993,118
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TYPE: amino acids
STRANDEDNESS: single stranded
TOPOLOGY: linear
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   NUMBER OF SEQUENCES: 14
CORRESPONDENCE ADDRESS:
ADDRESSER: Felfe & Lynch
STREET: 805 Third Avenue
CITY: New York City
STATE: New York
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INFORMATION FOR SEQ ID NO: 1
SEQUENCE CHARACTERISTICS:
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CURRENT APPLICATION DATA:
APPLICATION NUMBER: US
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US-08-993-118-10
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Sophie; DE SMET, Charles; BOON-FALLEUR, Thierry
: ISOLATED NUCLEIC ACID MOLECULE CODING
: FOR TUMOR REJECTION ANTIGEN PRECURSOR MAGE-C1 AND MAGE-C2
: AND USES THEREOF
                                                                                                                    190 DVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILLLSIIFIEGYCTPEEVIWEA 249
                                                                                                                                             70 TPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQVLPDSESL 129
                                                              250 LINMMGLYDGMEHLIYGEPRKLLITQDWVQENYLEYRQVPGSDPARYEFLWGPRAHAEIRKM 309
                                        130 PRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEASECMLLVFGI 189
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10 CMPEEDLOSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSSSSCYPLIPS 69
42 ILEEVPTAGST-DPPQSPQGASAFPTTINFTRQRQPSEGSSSREEEGPSTSCIL---ESL 97
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           - JEL/MAS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Diskette, 3.5 inch, 360 kb storage
                                                                                                                                                                                                                                                                                     310 SLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
                                                                                                                                                                                                                                                                                                                              278 KVLEYVIKVSARVRFFFPSLREAALREEEE 307
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       SOFTWARE: Wordperfect
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/066,281B
FILING DATE: April 24, 1998
CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION DATA:
APPLICATION NUMBER: 08/845,528
FILING DATE: April 25, 1997
ATTORNEY/AGENT INFORMATION:
NAMM: Mary Anne Schofield
REGISTRATION NUMBER: 36,669
REGISTRATION NUMBER: LUD 5455.2 U
TELEPHONE: (212) 318-3100
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Sequence 10, Application US/09066281B Patent No. 6475783 GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       os: single stranded
linear
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INFORMATION FOR SEQ ID NO: 1
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           APPLICANT: LUCAS, SOPHI
TITLE OF INVENTION: ISO
TITLE OF INVENTION: FOR
TITLE OF INVENTION: AND
NUMBER OF SEQUENCES: 20
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            COMPUTER READABLE FORM:
MEDIUM TYPE: Diskett
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  IBM PS/2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       TYPE: amino acids
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OPERATING SYSTEM:
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                                        DVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGYCTPEEVIWEA 249
                                                                      DVKEADPTGHSYVLVTCLGLSYDGLLGDNQIMPKTGFLIIVLVMIAMEGGHAPEREIWEE 217
                                                                                                                                                TPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQVLPDSESL 129
FRAVITKKVADLVGFLLLKYRAREPVTKAEMLESVIKNYKHCFPEIFGKASESLQLVFGI 157
                                                                                                                      LNNMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGPRAHAEIRKM 309
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MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFFWARE: Patentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/627,820
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
42.7%; Score 807; DB 4;
Best Local Similarity 53.6%; Pred. No. 7.6e-69;
Matches 177; Conservative 40; Mismatches 81
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Sidney, John C.
TITLE OF INVENTION: CLONING AND CHARACTER
COMPLETE MAGE 1 GENE
                                                                                                                                                                                                     310 SLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
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278 KVLEYVIKVSARVRFFFPSLREAALREEEE 307
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REGISTRATION NUMBER: 31,990
REGISTRATION INDER: 31,990
TELECOMMUNICATION INFORMATION:
TELEPHONE: (206) 467-9600
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         APPLICATION NUMBER: US/08/103,623
FILING DATE: <Unknown>
ATTORNEY/AGENT INFORMATION:
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MOLECULE TYPE: protein
SEQUENCE DESCRIPTION: SEQ ID NO: 24:
                                                                                                                                                                                                                                                                                                                                               Sequence 24, Application US/08627820
Patent No. 6464980
GENERAL INFORMATION:
APPLICANT: Fikes, John D.
Livingston, Brian D.
Sette, Alegandro D.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   FILING DATE: 02-Apr-1996
CLASSIFICATION: <Unknown>
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LENGTH: 309 amino acids
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        NUMBER OF
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DVKEADPTGHSYVLVTCLGLSYDGLLGDNQIMPKTGFLIVLVMIAMEGGHAPEEEIWEE 217
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                      TPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQVLPDSESL 129
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                                              130 PRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHPPLLPSEASECMLLVFGI
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APPLICANT: Scanlan, Matthew J.
APPLICANT: Gure, Ali O.
APPLICANT: Williamson, Barbara
APPLICANT: Chen, Yao-Teeng
APPLICANT: Chen, Yao-Teeng
APPLICANT: Old, Lloyd J.
ITILE OF INVENTION: Cancer Associated Antigens and Uses
ITILE OF INVENTION: Therefor
ITILE OF INVENTION: Therefor
TITLE OF ILLING DATE: 1999-09-09
CURRENT APPLICATION NUMBER: US/09/392,714A
CURRENT APPLICATION NUMBER: PCT/US98/14679
EARLIER FILING DATE: 1998-07-15
NUMBER OF SEQ ID NOS: 30
SEQ ID NO 29
LENGTH: 309
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42.7%; Score 807; DB 4;
Best Local Similarity 53.6%; Pred. No. 7.6e-69;
Matches 177; Conservative 40; Mismatches 81
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278 KVLEYVIKVSARVRFFPSLREAALREEBB 307
                                                                                                                                                                                                                                                                                                                               310 SLLKFLAKVNGSDPRSFPLWYBEALKDEEE 339
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                Sequence 29, Application US/09392714A Patent No. 6686147 GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ; TYPE: PRT
; ORGANISM: Homo sapiens
US-09-392-714-29
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APPLICANT: LUCAS, Sophie; BOON-FALLEUR, Thierry
TITLE OF INVENTION: ISOLATED NUCLEIC ACID MOLECULES CODING FOR
TITLE OF INVENTION: TUMOR REJECTION ANTIGEN PRECURSORS OF MEMBERS OF THE MAGE-C AN
TITLE OF INVENTION: MAGE-B FAMILIES AND USES THERROP
                                                                                                                                      DVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILLILSIIFIEGYCTPEEVIWEA 249
                                                                                                                                                                                           250 INMMGLYDGMEHLIYGEPRKLLITQDWVQENYLEYRQVPGSDPARYEFLWGPRAHABIRKM 309
                                                                                                                                                                                                                 130 PRSEIDEKVTDIVQFLLFKYQMKEPITKABILESVIKNYEDHFPLLFSEASECMLLVFGI 189
                                                          TLEEVPTAGST-DPPQSPQGASAFPTTINFTRQRQPSEGSSSREEEGPSTSCIL---ESL 97
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MEDIUM TYPE Diskette, 3.5 inch, 360 kb storage
COMPUTER: IBM PS/2
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/468,433C
FILING DATE: December 17, 1999
CLASSIFICATION NUMBER: US/09/468,231
FILING DATE: APPLICATION DATA:
APPLICATION NUMBER: 09/066,281
FILING DATE: APPTI 24, 1998
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/45,528
FILING DATE: APPTI 25, 1997
ATTORNEY/AGENT INFORMATION:
NAME: MATY ANDE SOLOFIELD
REFERENCE/DOCKET NUMBER: 1505
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10 CKPEEALEAQQEALGLVCVQA-----ATSSSS-----
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; Pred. No. 7.6e-69;
40; Mismatches 81
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278 KVLEYVIKVSARVRFFFPSLREAALREEEE 307
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   E: Fulbright & Jaworski L.L.P.
801 Pennsylvania Avenue, NW
                                                                                                                                                                                                                                                                                                                                                                                                      Sequence 10, Application US/09468433C Patent No. 6680191
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          TYPE: amino acids
STRANDEDNESS: single stranded
TOPOLOGY: linear
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       CITY: Washington
STATE: District of Columbia
COUNTRY: USA
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53.6%;
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INFORMATION FOR SEQ ID NO: 1
SEQUENCE CHARACTERISTICS:
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Best Local Similarity 53.6
Matches 177; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           NUMBER OF SEQUENCES: 2
CORRESPONDENCE ADDRESS:
ADDRESSEE: Fulbright
STREET: 801 Pennsylv
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                                                                                                                        FACULANT: VENTER, J. Craig et al.
APPLICANT: VENTER, J. Craig et al.
TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF
TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF
TITLE REPERENCE: CLOO1307
CURRENT APPLICATION NUMBER: US/09/949,016
CURRENT ELING DATE: 2000-04-14
PRIOR PLILING DATE: 2000-10-20
PRIOR FILING DATE: 2000-10-03
PRIOR FILING DATE: 2000-10-03
PRIOR FILING DATE: 2000-09-08
NUMBER OF SEQ ID NOS: 207012
SOUTHARD PRESEQ FOR WINDOWS VERSION 4.0
SEQ ID NO 6574
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Batent No. 6812339

GENERAL INFORMATION:
APPLICANT: VENTER, J. Craig et al.
ITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED
TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF
FILE REFERENCE: CLO01307

CURRENT APPLICATION NUMBER: US/09/949,016

CURRENT PILING DATE: 2000-04-14

PRIOR PILING DATE: 2000-10-20

PRIOR PILING DATE: 2000-10-20

PRIOR FILING DATE: 2000-10-03

PRIOR FILING DATE: 2000-10-03

PRIOR PLILNG DATE: 2000-10-03
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 309;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             40; Mismatches 81; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       42.7%; Score 807; DB 4; 53.6%; Pred. No. 7.6e-69;
KVLEYVIKVSARVRFFFPSLREAALREERE 307
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278 KVLEYVIKVSARVRFFFPSLREAALREEEE 307
                                                                                          Sequence 6574, Application US/09949016
Patent No. 6812339
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match 42.7%;
Best Local Similarity 53.6%;
Matches 177; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                              TYPE: PRT
ORGANISM: Human
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          US-09-949-016-8559
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Sequence 22, Application US/09468433C
Patent No. 6680191
GENERAL INFORMATION:
APPLICANT: LUCAS, Sophie; BOON-FALLEUR, Thierry
TITLE OF INVENTION: ISOLATED NUCLBIC ACID MOLECULES CODING FOR
TITLE OF INVENTION: TUMOR REJECTION ANTIGEN PRECURSORS OF MEMBERS OF THE MAGE-C ANT
TITLE OF INVENTION: MAGE-B FAMILIES AND USES THEREOF
NUMBER OF SEQUENCES: 26
CORRESPONDENCE ADDRESS:
ADDRESSEB: Pulbright & Jaworski L.L.P.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     190 DVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGYCTPEEVIWEA 249
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  204 DVKEADPTGHSYYLVTCLGLGYDGLLGDNQIMPKTGFLIIVLVMIAMEGGHAPBEEIWEE 263
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            70 TPEEVSADDETPNPPOSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQVLPDSESL 129
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             130 PRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKOYEDHFPLLFSEASECMLLVFGI 189
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                                                                                                                                                                                                                                                                                                                                               10 CMPREDIOSOSETOGLEGAQAPLAVEEDASSSTSTSSSFPSSSFPSSSSSSSSSCYPLIPS
                                                                                                                                                                                                                                                                                                                                                                                      CKPBEALEAQQSALGLVCVQA-----ATSSSS------PLVLG
                                                                                                                                                                                                                                      42.7%; Score 807; DB 4; Length 355; 53.6%; Pred. No. 9.6e-69;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette, 3.5 inch, 360 kb storage COMPUTER: IBM PS/2
OPERATING SYSTEM: PC-DOS
                                                                                                                                                                                                                                                                                                81;
                                                                                                                                                                                                                                                                                           40; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       310 SLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     :|:::||:
324 KVLEYVIKVSARVRFFFPSLREAALREEEE 353
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADDRESSER: Fulbright & Jaworski L.L.P. STREET: 801 Pennsylvania Avenue, NW CITY: Washington STATE: District of Columbia COUNTRY: USA
PRIOR FILING DATE: 2000-09-08
NUMBER OF SEQ ID NOS: 207012
SOFTWARE: FastSEQ for Windows Version 4.0
SEQ ID NO 8559
LENGTH: 355
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              SOFTWARE: Wordperfect
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/468,433C
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CLASSIFICATION
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 09/066,281
FILING DATE: April 24, 1998
FILING DATE: April 24, 1998
FILING DATE: April 25, 1997
ATTORNEY/ABGRY INCRMATION:
NAME: Mary Anne Schoffeld
REGISTRATION NUMBER: 36,669
                                                                                                                                                                                                                                                                                           Matches 177; Conservative
                                                                                                                                                                                                                                                                  Similarity
                                                                                                                                  ; TYPE: PRT
; ORGANISM: Human
US-09-949-016-8559
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US-09-468-433C-22
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 180 SECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILLILSIIFIEGY 239
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   240 CTPEEVIWEALNMMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWG 299
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             APPLICANT: Stroobant, Vincent
APPLICANT: Stroobant, Vincent
APPLICANT: Boon-Falleur, Thierry
APPLICANT: van der Bruggen, Pierre
APPLICANT: Thielemans, Kris
APPLICANT: Corthals, Jurgen
TITLE OF INVENTION: MAGE-3 PEPTIDES PRESENTED BY HLA CLASS II MOLECULES
FILE REPERRNCE: L0461/7052
CURRENT APPLICATION WUMBER: US/09/166,448
CURRENT PILING DATE: 1998-10-05
NUMBER OF SEQ ID NOS: 81
SOFTWARE: FastSEQ for Windows Version 3.0
SEQ ID NO 2
LENGTH: 314
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                155 SSSLQLVPGIELMEVDPIGHLYIFATCLGLSYDGLLGDNQIMPKAGLLIIVLAIIAREGD
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48.5%; Pred. No. 1.4e-66;
iive 59; Mismatches 89;
                                                                                                                                                                                                                                                                                                 DB 2;
                                                                                                                                                                                                                                                                                              Query Match 41.4%; Score 783.5; DB 2; Best Local Similarity 48.5%; Pred. No. 1.4e-66; Matches 165; Conservative 59; Mismatches 89;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Sequence 2, Application US/09166448 Patent No. 6291430
                                   INFORMATION FOR SEQ ID NO: 2: SEQUENCE CHARACTERISTICS: LENGTH: 314 amino acida TYPE: amino acida cinale
     TELECOMMUNICATION INFORMATION
                         : 617-720-3500
617-720-2441
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    GENERAL INFORMATION:
APPLICANT: Chaux, Pascal
APPLICANT: Vantomme, Valrie
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                                                                                                                                                                      STRANDEDNESS: single
TOPOLOGY: linear
                                                                                                                                                                                                                         , MOLECULE TYPE: protein US-08-928-615-2
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US-09-166-448-2
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          232 DLTYEGSLIDDQGMPKNCLLILILSMIPIKGSCVPEEVIWEVLSAIGVCAGREHFIYGDP 291
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     88 QIACSSPSVVASLPLDQSDEGSSSQKEESPSTLQVLPDSESLPRSEIDEKVTDLVQFLLF 147
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                                                                                                                                                                                                                                                                                                                                                                                                                                 1 MPLFPNLPRLSFEEDPQNPSVTEDLVDAQDSIDEEEEDASSTS-SSSFHFLFPSSSSLSS
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    RKLLTQDWVQENYLEYRQVPGSDPARYEFLWGPRAHAEIRKMSLLKFLAKV 318
                                                                                                                                                                                                                                                                                                                                                    42;
                                                                                                                                                                                                                                                                                                    DB 4; Length 346;
                                                                                                                                                                                                                                                                                                 Query Match
41.9%; Score 793; DB 4; Length 346
Best Local Similarity 49.9%; Pred. No. 2e-67;
Matches 175; Conservative 43; Mismatches 91; Indels
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US-08-928-615-2
Sequence 2, Application US/08928615
Sequence 2, Application US/08928615
Sequence 2, Application US/08928615
Sequence 3, Application US/08928615
Sequence 4, Paccal APPLICANT Chaux, Pascal APPLICANT Broom, Thierry
APPLICANT Boon, Thierry
APPLICANT To an dear Bruggen, Pierre APPLICANT TO FINVENTION: BY HLA CLASS II MOLECULES
INMERS OF SEQUENCES: 13
CORRESPONDENCE ADDRESS:
ADDRESSEE: Wolf, Greenfield & Sacks, P.C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            OPERATING SYSTEM: DOS
SOFTWARE: FastSEQ for Windows Version 2.0
CHRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/928,615
REFERENCE/DOCKET NUMBER: LUD 5611 JEL/MAS
TELECOMMUNICATION INFORMATION:
TELEPHONE: (202) 662-0200
TELEFAX: (202) 662-4643
INFORMATION FOR SEQ ID NO: 22:
SEQUENCE CHARACTERISTICS:
LENGTH: 346
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             E: Wolf, Greenfield & Sacks, P.C.
600 Atlantic Avenue
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      61 SSCYPL---IPST------
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NAME: Van Amsterdam, John R.
REGISTRATION NUMBER: 40,212
REFERENCE/DOCKET NUMBER: L0461/7017
                                                                                                                                                                         TYPE: amino acid
STRANDEDNESS: single-stranded
TOPOLOGY: linear
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            COUNTRY: U.S.A.
ZIP: 02210-2211
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: IBM COMPATIBLE
OPERATING SYSTEM: DOS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       CLASSIFICATION: 424
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                                                                                                                                                                                                                            ; TOPOLOGY:
US-09-468-433C-22
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APPLICANT: Chaux, Pascal
APPLICANT: Chaux, Pascal
APPLICANT: Chaux, Vincent
APPLICANT: Boon-Falleur, Vincent
APPLICANT: Wonder Bruggen, Pierre
TITLE OF INVENTION: MAGE-3 PEPTIDES PRESENTED BY HLA CLASS II MOLECULES
FILE REFERENCE: LO461/7065
CURRENT APPLICATION NUMBER: US/09/348,933
CURRENT FILING DATE: 1999-07-07
EARLIER APPLICATION NUMBER: US 08/928,615
BARLIER PILING DATE: 1997-09-12
NUMBER OF SEQ ID NOS: 13
SOFTWARE: RestSEQ for Windows Version 3.0
SEQ ID NO 2
LENGTH: 214
                                                                                                                                               180 SECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILILSIIFIEGY 239
                                                                                                                                                                    CTPERVIWEALNAMGLYDGMEHLIYGEPRKILITQDWVQENYLEYRQVPGSDPARYEFLWG 299
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     61 SSCYPLIPSTPEEVSADDETPNPPOSAQIACSSPSVVASLPL-DQSDEGSSSQKEESPST 119
61 SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPL-DQSDEGSSSQKEESPST 119
                                                                                                                                                                                                                                             215 CAPEEKIWEELSVLEVFEGREDSILGDPKKLITQHFVQENYLEYRQVPGSDPACYEFLWG 274
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               120 LQVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKBPITKABILESVIKNYEDHPPLLFSEA 179
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      98 ---FPDLESEFQAALSRKVAELVHFILLKYRAREPVTKAEMLGSVVGNWQYFFPVIFSKA 154
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                     45 -----LVEVTLGEVPA-AESPDPPQSPQGASSLPTTM-NYPLWSQSYEDSSNQEEEGPSI 97
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Patent No. 6369211
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ORGANISM: Homo sapiens
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Best Local Similarity
Matches 165; Conserval
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Search completed: November 15, 2005, 15:45:52 Job time : 62.5476 secs

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Sequence 32058, A
Sequence 188, App
Sequence 52, Appli
Sequence 148, App
Sequence 2, Appli
Sequence 2, Appli
Sequence 40, Appli
                                                                                 November 15, 2005, 15:35:58; Search time 219.643 Seconds (without alignments) 702.928 Million cell updates/sec
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score greater than or equal to the score of the result being printed,
and is derived by analysis of the total score distribution.
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GenCore version 5.1.6
(c) 1993 - 2005 Compugen Ltd.
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US-10-188-832-149
US-10-658-884-4
US-10-756-149-4721
US-10-029-386-32058
US-10-482-029-188
US-10-157-031-52
US-10-75-149-4740
US-10-75-18-095-2
US-10-093-766-40
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Maximum Match 100%
Listing first 45 summaries
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Gapop 10.0 , Gapext 0.5
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Maximum DB seq length: 200000000
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        US-10-157-031-54
        Sequence 54, Appl 466.5
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        US-10-473-127-1069
        Sequence 1079, Appl 56.4
        Appl 56.5
        44.8
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        US-10-473-127-1029
        Sequence 1091, Appl 66.5
        Appl 56.5
        44.8
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        US-10-473-127-1029
        Sequence 1091, Appl 76.5
        Appl 76.5
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        US-10-473-127-1029
        Sequence 1122, Appl 76.4
        Appl 76.5
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        US-10-473-127-1029
        Sequence 1122, Appl 76.4
        Appl 76.5
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        US-10-473-127-1029
        Sequence 1122, Appl 76.4
        Appl 76.7
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ALIGNMENTS

RESULT 1
US-10-05-542-84

| Sequence 84, Application US/10036542
| Sequence 84, Application US/10036542
| Publication No. US20030083481A1
| GRNERL INFORMATION:
| APPLICAMTI SITES et al. |
| TILLE OF INVENTION: 2 Human Prostate and Prostate Cancer Associated Proteins
| TILLE OF INVENTION: 2 Human Prostate and Prostate Cancer Associated Proteins
| TILLE OF INVENTION: 2 1002-01-07
| PRIOR PILLING DATE: 2000-07-20
| PRIOR FILLING DATE: 1999-07-21
| PRIOR FILLING DATE: 1999-07-21
| PRIOR FILLING DATE: 1999-08-17
| PRIOR FILLING DATE: 1999-08-17
| PRIOR APPLICATION NUMBER: 60/144,972
| PRIOR APPLICATION NUMBER: 60/149,173
| PRIOR APPLICATION NUMBER: 60/149,173
| PRIOR APPLICATION NUMBER: 60/158,004
| PRIOR FILLING DATE: 1999-10-06
| PRIOR FILLING DATE: 2000-04-05
| NUMBER OF SEQ ID NOS: 157
| ORGANISM: Home sapiens
| US-10-036-542-84
| Query Match
| DROWN MATCH MATC

240

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QVLPDSSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS 180
                                                                                                                                                                                                          TPEEVIWEALNAMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVPGSDPARYEFLWGP 300
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                                                                                                                                                                                                                                                                                                                RAHAEIRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEBERAQDRIATTDDTTAMASASSS 360
QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              TITLE OF INVENTION: NOVEL HUMAN MAGE-LIKE PROTEIN
NUMBER OF SEQUENCES: 6
CORRESPONDENCE ADDRESS:
ADDRESSER: Incyte Pharmaceuticals, Inc.
STREET: 3174 Porter Drive
CITY: Palo Alto
STATE: CA
COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        COMPUTER READABLE FORM:

MEDIUM TYPE: DISKETE
COMPUTER: IBM Compatible
COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FASTENG for Windows Version 2.0
CURRIT APPLICATION DATA:
APPLICATION NUMBER: US/10/658,884
FILING DATE: 09-8ep-2003
CLASSIPTICATION 435
PRIOR APPLICATION A435
PRIOR DATE: 37-BEC-1996
ATTORNYA/AGENT INFORMATION:
NAME: Billings, Lucy J.
REGISTRATION NUMBER: 36,749
REFERENCE/DOCKET NUMBER: 36,749
TELECOMMUNICATION INFORMATION:
TELECOMMUNICATION INFORMATION:
TELECOMMUNICATION INFORMATION:
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SEQUENCE DESCRIPTION: SEQ ID NO: 4:
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TELEX: 40htnonn>
INFORMATION FOR SEQ ID NO: 4:
SEQUENCE CHARACTERISTICS:
LENGTH: 369 amino acide
TYPE: amino acide
STRANDEDNESS: single
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Publication No. US20040076955A1
GENERAL INPORMATION:
APPLICANT: Mack, David H.
APPLICANT: Mack, David H.
APPLICANT: AZIZ, Natasha
TITLE OF INVENTION: Methods of Diagnosis of Bladder
TITLE OF INVENTION: Methods of Screening for Modulators of Bladder
TITLE OF INVENTION: and Methods of Screening for Modulators of Bladder
TITLE OF INVENTION: Cancer
FILE REFERENCE: 018691-002330US
CURRENT APPLICATION NUMBER: US/10/188,832
CURRENT APPLICATION NUMBER: US 60/3302,814
PRIOR FILING DATE: 2001-07-03
PRIOR FILING DATE: 2001-07-03
PRIOR FILING DATE: 2001-11-13
PRIOR FILING DATE: 2001-11-13
PRIOR APPLICATION NUMBER: US 60/343,705
PRIOR APPLICATION NUMBER: US 60/372,246
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                                                                                                  SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL 120
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                           1 MPRAPKRQRCMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSS
WPRAPKRORCMPEEDLOSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSS
                                                                                                                                      SSCYPLIPSTPEEVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL
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100.0%; Pred. No. 3.1e-141;
ive 0; Mismatches 0;
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US-10-188-832-149
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Qy 301 RAHAEIRKMSLLKFLAKVNGSDPRGPPLWYBEALKDEBERAQDRIATTDDTTAMAGASS.	RESULT 5 US-010-029-386-32058 US-010-029-386-32058 Sequence 32058, Application US/10029386 Publication No. US20030194704A1 GENERAL INFORMATION: APPLICANT: Penn, Sharron G. APPLICANT: HAnzel, David R. TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE CACID PROBES TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE STON NUCLEIC ACID PROBES CURRENT APPLICATION NUMBER: US/10/029,386 CURRENT PILING DATE: 2001-12-20 NUMBER OF SEQ ID NOS: 34288 SOFTWARE: Annomax Sequence Listing Engine vers. 1.1 SEQ ID NO 32058 LENGTH: 383 TYPE: PRT	CHER INFORMATION: MAP TO AF134576.1 CTHER INFORMATION: EXPRESSED IN ADULT LIVER, SIGNAL = 1.7 OTHER INFORMATION: EXPRESSED IN LONG, SIGNAL = 4.1 OTHER INFORMATION: EXPRESSED IN PLACENTA, SIGNAL = 1.4 OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 1.6 OTHER INFORMATION: EXPRESSED IN HEART, SIGNAL = 1.6 OTHER INFORMATION: EXPRESSED IN HEART, SIGNAL = 1.9 OTHER INFORMATION: EXPRESSED IN HEART, SIGNAL = 1.9 OTHER INFORMATION: EXPRESSED IN BONE MARROW, SIGNAL = 1.8 OTHER INFORMATION: EXPRESSED IN BONE MARROW, SIGNAL = 1.8 OTHER INFORMATION: EXPRESSED IN FORMATION = 1.6 OTHER INFORMATION: EXPRESSED IN FORMATION = 0.00e+00 US-10-029-386-32058 QUARTY MATCH PRAPKRORCMPEEDLOSOSETOGLEGAQAPLANEEDASSESTESSSPPSSFPSSSSSSS DD 22 MARAPKRORCMPEEDLOSOSETOGLEGAQAPLANEEDASSESTESSSFPSSFPSSSSSSSSDD DD 22 MRAAPKRORCMPEEDLOSOSETOGLEGAQAPLANEEDASSSTSTSSSFPSSFPSSSSSSSSDD	202 202 301 301 301 301 301
MPRAPKRQRCMPEEDLOSQSETGGLEGAQAPLAVEEDASSSTSTSSSFPSSSSSS 60	121 QVLPDSESLPRSEIDEKVTDLVQFLLFKXQMKEPITKAEILESVIKNYEDHFPLLFSEAS 180 121 QVLPDSESLPRSEIDEKVTDLVQFLLFKXQMKEPITKAEILESVIKNYEDHFPLLFSEAS 180 121 QVLPDSESLPRSEIDEKVTDLVQFLLFKXQMKEPITKAEILESVIKNYEDHFPLLFSEAS 180 181 ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILLILSIIFIEGYC 240 181 ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILLILSIIFIEGYC 240 181 ECMLLVFGIDVKEVDPTGHSFVLVTSLGLTYDGMLSDVQSMPKTGILILLILSIIFIEGYC 240 241 TPEEVIWEALNYMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVFGSDPARYEFLWGP 300 241 TPEEVIWEALNYMGLYDGMEHLIYGEPRKLLTQDWVQENYLEYRQVFGSDPARYEFLWGP 300 301 RAHAEIRKMSLLKFLAKVNGSDPRSFPLMYEEALKDEEERAQDRIATTDDTTAMASASS 360 1	RESULT 4 US-10-756-149-4721 Sequence 4721, Application US/10756149 Publication No. US20050181375A1 GENERAL INFORMATION: APPLICANT: Aziz, Natasha APPLICANT: Aziz, Natasha TITLE OF INVENTION: NOVEL METHODS OF DIAGNOSIS OF METASTATIC CANCER, TITLE OF INVENTION: METHODS OF SCREENING FOR MODULATORS OF METASTATIC CANCER TITLE OF INVENTION: MATHODS OF SCREENING FOR MODULATORS OF METASTATIC CANCER TITLE OF INVENTION: MATHODS OF SCREENING FOR MODULATORS OF METASTATIC CANCER CURRENT FILLING DATE: 2004-01-12 NUMBER OF SEQ ID NOS: 5818 SEQ ID NO 4721 LENGTH: 369 TYBE: PRT TYBE: PRT TYBE: PRT TYBE: PRT CREATION: MATHOD SADIENS CREATION: MATHOD SADIENS CORRANS FOR METASTATIC CANCER CORRANS FOR METASTATIC CANCER TOTAL FOR METASTATIC CANCER TITLE OF METASTATIC CANCER	Scheme Section Secti

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                                                                               61 SSCYPLIPSTPREVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL
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                           1 MSSEQKSQHCKPEEGVEAQEEALGLVGAQAPTTEEQEAAVSSSS-
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Best Local Similarity 51.9%; Pred. No. 3.7e-59;
Matches 176; Conservative 51; Mismatches 88;
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; Bedication No. US20050037445A1
; GENERAL INFORMATION:
; APPLICANT: ODIN medical A/S
; TITLE OF INVENTION: Oncology drug innovation
; FILE REFERENCE: P 573 PC00
; CURRENT APPLICATION NUMBER: US/10/482,029
; CURRENT FILING DATE: 2003-12-29
; NUMBER OF SEQ ID NOS: 437
; SOFTWARE: Patentin version 3.1
; SEQ ID NO 148
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CRGANISM: Homo sapiens
US-10-482-029-148
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APPLICANT: Yankovsky, N. K.
APPLICANT: Yozlov, A. P.
APPLICANT: Kozlov, A. P.
APPLICANT: Kozlov, A. P.
APPLICANT: Kozlov, A. V.
APPLICANT: Kolovoskaya, L. L.
TITLE OF INVENTION: In silico screening for phenotype-associated expressed sequences
FILE REFERENCE: 2760-103
CURRENT APPLICATION NUMBER: US/10/157,031
CURRENT FILING DATE: 2002-05-30
NUMBER OF SEQ ID NOS: 415
SOFTWARE: Patentin version 3.1
SEQ ID NO 52
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                                                                                                                                                                                                                                                                                                                                                               48.2%; Score 911; DB 17; Length 319; 54.0%; Pred. No. 1.1e-63; ive 53; Mismatches 81; Indels 2;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ; Sequence 52, Application US/10157031; Publication No. US20030108890A1; GENERAL INFORMATION:
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; ORGANISM: Homo sapiens
US-10-482-029-188
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; ORGANISM: Homo sapiens
US-10-157-031-52
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US-10-157-031-52
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Best Local
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Publication No. US20030013099A1

GENERAL INFORMATION:

APPLICANT: Lasek, Amy W.

APPLICANT: Karpf, Adam R.

TITLE OF INVENTION: GENES REGULATED BY DNA METHYLATION IN COLON TUMORS

FILE REFERENCE: PA-0047 US/10/093,766

CURRENT PILING DATE: 2002-03-07

NUMBER OF SEQ ID NOS: 61

SEQ ID NO 40

LENGTH: 315
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    MPRAPKRORCMPEEDLOSOSETOGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSS
                                                                                                                    45 ---- PLVPGTLEEVPA-AESAGPPQSPQGASALPTTISFTCWRQPNEGSSSQEEEGPST-
                                                                              61 SSCYPLIPSTPERVSADDETPNPPQSAQIACSSPSVVASLPLDQSDEGSSSQKEESPSTL
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OTHER INFORMATION: Incyte ID No. US20030013099A1 2502336CD1
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Matches 172; Conservative
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US-10-093-766-40
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Sequence 4740, Application US/10756149
Publication No. US20050181375A1
GENERAL INFORMATION:
APPLICANT: ALLA, Natasha
APPLICANT: Zlotnik, Albert
APPLICANTON: METHODS OF DIAGNOSIS OF METASTATIC CANCER, COMPOSITIONS AND TITLE OF INVENTION: METHODS OF SCREENING FOR MODULATORS OF METASTATIC CANCER OURRENT PILING DATE: 2004-01-12
NUMBER OF SEQ ID NOS: 5818
SOFTWARE: Patentin version 3.2
SEQ ID NO 4740
LENGTH: 217
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             121 QVLPDSESLPRSEIDEKVTDLVQFLLFKYQMKEPITKAEILESVIKNYEDHFPLLFSEAS
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; GENERAL INFORMATION:
APPLICANT: Stroobart, Vincent
APPLICANT: Boon-Falleur, Thierry
APPLICANT: US BOON-Falleur, Thierry
APPLICANT: US BOON-Falleur, Thierry
APPLICANT: US BOON-Falleur, Thierry
APPLICANT: US BOON-Falleur, Thierry
CURRENT APPLICATION NUMBER: US/10/218,095
CURRENT APPLICATION NUMBER: US/10/218,095
CURRENT PILING DATE: 2002-08-13
NUMBER OF SEQ ID NOS: 63
; SEQ ID NO 2
LENGTH: 317
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 317;
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;; Pred. No. 3.7e-59;
51; Mismatches 88; Indels 2
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51.6%; Pred. No. 9.2e-59;
ive 51; Mismatches 89; Indels
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Best Local Similarity 51.9%;
Matches 176; Conservative 5:
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Best Local Similarity 51.6
Matches 175; Conservative
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ORGANISM: Homo sapiens
US-10-218-095-2
                                                                                                                                                                                                                                                                                    TYPE: PRT
ORGANISM: Homo Sapiens
                                                                                                                                                                                                                                                                                                                         US-10-756-149-4740
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97 PAQLEFWFQEALKLKVAELVHFILHKYRVKBPVTKAEMLESVIKNYKRYFPVIFGKASEF 156
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                  CURRENT APPLICATION NUMBER: US 1010/10/295,027

CURRENT PAPLICATION NUMBER: US 109/663,733

PRICA APPLICATION NUMBER: US 09/663,733

PRICA APPLICATION NUMBER: US 09/663,733

PRICA PILING DATE: 2000-09-15

PRICA PILING DATE: 2001-11-13

PRICA PILING DATE: 2001-11-12

PRICA PILING DATE: 2001-11-12

PRICA PILING DATE: 2001-11-29

PRICA PILING DATE: 2001-12-14

PRICA PILING DATE: 2001-12-14

PRICA PILING DATE: 2001-12-14

PRICA PILING DATE: 2002-01-08

PRICA PILING DATE: 2002-01-13

PRICA PILING DATE: 2002-01-13

PRICA PILING DATE: 2002-01-13

PRICA PILING DATE: 2002-01-13

PRICA PILING DATE: 2002-02-13

PRICA PILING DATE: 2002-03-13

PRICA PILING DATE: 2002-03-13
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Publication No. US20040236091A1
GENERAL INFORMATION:
APPLICANT: Zycos Inc.
TITLE OF INVENTION: TRANSLATIONAL PROFILING
FILE REFERENCE: 08191-026801
CURRENT APPLICATION NUMBER: US/10/473,127
CURRENT FILING DATE: 2003-09-26
PRIOR PILING DATE: 2003-09-26
PRIOR PILING DATE: 2001-03-28
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US-10-295-027-366
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Best Local Similarity
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US-10-157-031-54

i Sequence 54, Application US/10157031

i Publication No. US20030108890A1

i Publication No. US20030108890A1

i GENERAL INFORMATION:

APPLICANT: Baranova, A. V.

APPLICANT: Kozlov, A. P.

APPLICANT: Krakovskaya, L. L.

TITLE OF INVENTION: In silico screening for phenotype-associated expressed sequences

FILE REPERENCE: 2760-103

CURRENT APPLICATION NUMBER: US/10/157,031

CURRENT APPLICATION NUMBER: US/10/157,031

CURRENT APPLICATION NUMBER: US/10/157,031

CURRENT PILING DATE: 2002-05-30

NUMBER OF SEQ ID NOS: 415

SSOTTWARE: Patentin version 3.1

LENGTH: 315
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               3 RAPKRORCMPEEDLQSQSETQGLEGAQAPLAVEEDASSSTSTSSSFPSSFPSSSSSSSSSSSS
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APPLICANT: Watson, Susan R.
APPLICANT: EOS Biotechnology, Inc.
TITLE OF INVENTION: Methods of Diagnosis of Cancer, Compositions and
TITLE OF INVENTION: Methods of Screening for Modulators of Cancer
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DB 14; Length 315;
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44.8%; Score 846.5; DB 14; Length
Best Local Similarity 51.0%; Pred. No. 1.4e-58;
Matches 172; Conservative 48; Mismatches 88; Indels
303 HAEIRKMSLLKFLAKVNGSDPRSFPLWYEEALKDEEE 339
                                           277 HAETSYEKVINYLVMLNAREPICYPSLYEEVLGEEQE 313
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Publication No. US20030232350Al
GENERAL INFORMATION:
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APPLICANT: Aziz, Natasha
APPLICANT: Giaberg, Wendy M.
APPLICANT: Gish, Kurt C.
APPLICANT: Glynne, Richard
APPLICANT: Hevezi, Peter A.
APPLICANT: Mack, David H.
APPLICANT: Marray, Richard
APPLICANT: Watson, Susan R.
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ORGANISM: Homo sapiens
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US-10-295-027-366
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US-10-157-031-54
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; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-473-127-1087
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                                                                                                                                                                                                                                                                                                                                                                                   44.8%; Score 846.5; DB 16; Length 315; 51.0%; Pred. No. 1.4e-58; tive 48; Mismatches 88; Indels 29;
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TITLE OF INVENTION:
FILE REPERENCE:
CURRENT ZYCOS INC.
TITLE OF INVENTION: TRANSLATIONAL PROFILING
FILE REPERENCE:
CURRENT APPLICATION NUMBER: US/10/473,127
CURRENT FILING DATE: 2003-09-26
FRIOR PELION NUMBER: 60/29,495
FRIOR FILING DATE: 2001-05-21
FRIOR PELION NUMBER: 60/292,544
FRIOR PILING DATE: 2001-05-01
FRIOR APPLICATION NUMBER: 60/30,00
FRIOR PILING DATE: 2001-06-08
FRIOR PILING DATE: 2001-06-01
FRIOR PILING DATE: 2001-06-01
FRIOR PILING DATE: 2001-10-01
FRIOR PILING DATE: 2001-10-01
FRIOR PILING DATE: 2001-10-01
FRIOR APPLICATION NUMBER: 60/336,780
FRIOR PILING DATE: 2001-12-04
FRIOR PILING DATE: 2001-12-04
FRIOR PILING DATE: 2001-12-04
FRIOR PILING DATE: 2002-02-20
FRIOR FILING DATE: 2002-02-20
FRIOR FILING DATE: 2003-03-20
FRIANES FABELSEQ FOR WINDOWS VERSION 4.0
FROM THE STATES FABELSED FOR WINDOWS VERSION 4.0
               PRIOR FILING DATE: 2001-05-21
PRIOR APPLICATION NUMBER: 60/310,801
PRIOR PILING DATE: 2001-08-08
PRIOR PLING DATE: 2001-08-08
PRIOR PLING DATE: 2001-10-01
PRIOR PILING DATE: 2001-10-01
PRIOR PILING DATE: 2001-12-04
PRIOR PILING DATE: 2002-02-20
NUMBER OF SEQ ID NOS: 2041
SEQ ID NO 1079
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Publication No. US20040236091A1
GENERAL INFORMATION:
  APPLICATION NUMBER: 60/292,544
                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity 51.0°
Matches 172; Conservative
                                                                                                                                                                                                                                                                                                 TYPE: PRT
ORGANISM: Homo sapiens
US-10-473-127-1079
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  DB 16; Length 315;
                                                      Indels
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Query Match
44.8%; Score 846.5; DB 16;
Best Local Similarity 51.0%; Pred. No. 1.4e-58;
Matches 172; Conservative 48; Mismatches 88;
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Search completed: November 15, 2005, 15:49:48 Job time : 227.643 secs

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November 15, 2005, 15:28:51; Search time 5.69048 Seconds (without alignments) 611.696 Million cell updates/sec
GenCore version 5.1.6 (c) 1993 - 2005 Compugen Ltd.
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                                                             - protein search, using sw model
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51
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              Copyright
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                                                               protein
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Perfect
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Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Total number of hits satisfying chosen parameters:

seq length: 0 seq length: 200000000

Minimum DB Maximum DB

geneseqp1980s:* geneseqp1990s:* geneseqp2000s:* geneseqp2001s:* geneseqp2002s:* geneseqp2003as:*geneseqp2003bs:*geneseqp2004s:* A_Geneseq_16Dec04:* Database

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

	Description	Aay71487 Human MAG	Aab31323 Exemplary	80		Adg89586 Class I H	_		Adj58374 Peptide p	0	Adr69760 Novel hyb	Aay71489 Human MAG		5 Human	7 Human	Abr48215 Human bla	Abu56516 Lung canc		Ada55634 Human pro	Aay71488 Human MAG	Aay71490 Human MAG	Aab08734 Amino aci	Abm82457 Tumour-as	Aaw24252 Ammonifex	Abu57355 A. degens	
SUMMAKIES	ΙD	AAY71487	AAB31323	AA017088	ABJ19876	ADG89586	ADI19869	ADI19026	ADJ58374	ADL26560	ADR69760	AAY71489	ADR69766	AAY71485	AAB80297	ABR48215	ABUS6516	AB058424	ADA55634	AAY71488	AAY71490	AAB08734	ABM82457	AAW24252	ABU57355	
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	Score	51	51	51	51	51	51	51	51	51	51	51	51	51	51	51	51	51	43	41	41	41	41	40	40	:
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Adj6989 Human hea Adn20149 Bacterial Abo79916 Bacterial Aby2270 MAGB-A4 a Aab11320 Exemplary Aac17089 Human mag Adj19875 MHC bindi Adj19874 Human HLA Adj19874 Human HLA Adj19874 Human HLA Adj5338 Peptide p Ado5916 Human MAG Ado5916 Human MAG Ado5916 Human MAG Ado5916 Peptide # Abb3645 Peptide # Abb3642 Peptide # Abb3082 Peptide # Abb3082 Peptide # Abb20872 Proteiln # Amm68635 Human bon	Aam56259 Human bra
ADJ69889 ADV20149 ABV7916 AAV792270 AAB31320 AAA119875 ADG19875 ADG19874 ADG19874 ADG19031 ADG59166 AAM1644 AAM1644 AAM1644 AAM1644 AAM169635 AAM28942 AAM306635	AAM56259
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	38
И С С С С С С С С С С С С С С С С С С С	45

ALIGNMENTS

MAGB-A10; human; Tumour Rejection Antigen; TRA; Human Leucocyte Antigen; HLA; Major Histocompatibility Complex; MHC; cytolytic T-lymphocyte; CTL; immune response stimulator; prophylaxis; therapy; diagnosis; tumour; cancer; TNF; tumour necrosis factor; vaccine; cytostatic. Boon T, De Plaen E, Brasseur F, AAY71487 standard; peptide; 9 AA. (LUDW-) LUDWIG INST CANCER RES. Human MAGE-Al0 nonapeptide-1. 98GB-00026143. 99WO-IB002018. (first entry) Van Pel A, WPI; 2000-412317/35. WO200032769-A2 26-NOV-1999; 27-NOV-1998; Homo sapiens 08-JUN-2000. 12-OCT-2000 AAY71487; Huang L,

Novel polypeptides expressed in tumor cells useful for treating cancers have an ability to complex with a major histocompatibility complex molecule and comprises a specific unbroken amino acid sequence.

Claim 8; Page 36; 80pp; English.

The patent discloses MAGE-A10 and WAGE-A8 polypeptide, nonapeptide and decapeptide sequences, that function as tumour rejection antigens (TRAs). These peptides are capable of forming a complex with major histocompatibility complex (MHC) molecule type HLA-A2.1 (Human Leucocyte Antigen), that are recognised by T-lymphocytes and elicit an immune response from cyclytic T-lymphocytes (TL). They function as an immune response stimulator. Tumour rejection antigens are useful in prophylaxis, therapy and diagnosis of tumours and are effective in controlling or preventing tumour growth. The present peptide sequence is the human MAGE-A10 nonapeptide-1, that corresponds to residues 254-262 of the MAGE-A10 protein. This peptide can serve as a tumour rejection antigen (TRA) and

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Sequence 9 AA;
                                                                                                                                                     Homo sapiens
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                                              AA017088;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AAB31302-59 represent exemplary antigens which are characteristic of tumours. They can be used to enhance the immune response of vaccines comprising peptides drived from human MAGE-A1 HLA (human leukocyte antigen) class II-bindring protein. Peptides derived from the MAGE-A1 HLA bindring protein stimulate the activity and proliferation of CD4+ T lymphocytes. The MAGE-A1 HLA bindring protein is useful as a disapnestic agent for diagnosing a disorder characterized by expression of MAGE-A1 such as cancers e.g. melanomal, squamous cell carcinomas, colorectal carcinomas, osteosarcomas, and lymphocytic leukemias. Peptides derived from the MAGE-A1 HLA bindring protein are useful in the production
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in combination with adjuvants, can produce vaccines useful for treating a variety of tumours that express MAGE-Al0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Novel MACE-A1 human leukocyte antigen class II peptides which bind to are presented to the class II molecules, useful for inducing immune response and treating cancers characterized by expression of MAGE-A1.
                                                                                                                                                                                                                                                          Exemplary antigen characteristic of tumours and derived from MAGE-AlO.
                                                                                                                                                                                                                                                                                                                                                                                                                                                            Van Der Bruggen P;
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                                                          Length 9;
                                                                               0; Indels
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                                                         Score 51; DB 3; I
Pred. No. 1.8e+06;
                                                                                Mismatches
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                                                                                                                                                                                     AAB31323 standard; peptide; 9 AA
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                                                         100.0%;
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                                             Query Match
Best Local Similarity 100.
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                                                                                                      1 GLYDGMEHL
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Sequence 9 AA;
                                   Sequence 9 AA;
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                                                                                                                                                                                                                                   20-APR-2001
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Antirheumatic; antiallergic; antiarthritic; nootropic; neuroprotective; antinflammatory; major histocompatibility complex; MHC; autoimmune disease; T cell; B cell; allergic disease; multiple sclerosis; rheumatoid arthritis; neurodegenerative disorder; Alzhelmer's disease; inflammation; gene therapy; MHC binding peptide.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                The present invention relates to a method for the preparation of readyfor-use, cryopreserved, mature dendritic cells comprising growing mature dendritic cells in a culture medium that includes a 'maturation cockeail' of one or more maturation stimuli and freezing the resulting matured cells in a freezing medium that does not contain heterologous everum. When loaded with antigens, the dendritic cells can be used as vaccines, e.g. against tumours and hepatitis B virus. The present sequence is an antigen described in the invention
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Preparation of cryopreserved, mature dendritic cells, useful in vaccines, comprises culturing immature cells on medium containing cocktail of
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                                                                                                                                                                                                   Cryopreserved mature dendritic cell; antigen; vaccine; cytostatic; virucide; cancer; hepatitis B virus.
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                                                                                                                                                       Human mage-A10 protein antigen SEQ ID NO: 8.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        maturation factors, then freezing.
AA017088 standard; peptide; 9 AA.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      24-AUG-2000; 2000DE-01041515.
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                                                                                                                                                                                                                                                                                                                                            WO200216560-A1.
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WPI; 2004-035134/03
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                                                                                                                                                                                                                                                                                 RESULT 6
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                                                                                                                                                                                                                                         The invention relates to a novel method for identifying peptides originating from a particular cell type, which are capable of binding to major histocompatibility complex (MHC) molecules of a particular haplotype. The method comprises analysing peptides bound to the soluble and secreted form of the MHC molecules of the particular haplotype. The method is useful for identifying peptides for treating an autoimmune disease, such as T or B cell and/or allergic disease or condition, rheumatoid arthritis, or multiple sclerosis, neurodegenerative disorders, e.g. Alzheimer's disease, or diseases associated with inflammation. The sequences of the invention may be used in a gene therapy application. This sequence represents a peptide relating to the method for identifying MHC binding peptides of the invention
                                                                                                                                                                 Identifying peptides that are capable of binding to major histocompatibility complex (WHC) molecules of a particular haplotype by analyzing peptides bound to the soluble and secreted form of the MHC molecules of the particular haplotype.
                                                                                                                                                                                                                                                                                                                                                                                                                                           Gaps
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                                                                                                                         Buchsbaum S;
                                                                                                                                                                                                                                                                                                                                                                                                                     Length 9;
                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match 100.0%; Score 51; DB 6; Length 9; Best Local Similarity: 100.0%; Pred. No. 1.8e+06; Matches 9; Conservative 0; Mismatches 0; Indels
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                                                                                                                         Давваи L,
                                                                                                                       Admon A,
                                                                                               (TECR ) TECHNION RES & DEV FOUND LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADG89586 standard; peptide; 9 AA.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (BAYU ) BAYLOR COLLEGE MEDICINE.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        28-MAY-2003; 2003WO-US016736.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              28-MAY-2002; 2002US-0383530P.
                                           16-MAY-2002; 2002WO-IL000383.
                                                               16-MAY-2001; 2001US-0290958P.
29-MAY-2001; 2001US-00865548.
                                                                                                                      Beer I, Ziv T,
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WO200294981-A2
                                                                                                                                                                                                                                                                                                                                                                                               Sequence 9 AA;
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                                                                                                                       Barnea E,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADG89586;
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8XX4X8X8XX1X8XX1X1X8XX1XX8XX0XXX
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Identifying a cell that differentiates into a metastatic cancer cell, useful for preventing metastatic cancer, comprises identifying a mutated fibronectin in the cell.
                                                                                                                                                                                                   The invention comprises a method for identifying a cell that will differentiate into a metastatic cancer cell, the method involves identifying a mutated fibronectin in the cell. The method of the invention is useful for preventing metastatic cancer. The present amino acid sequence represents a Class I HLA-restricted testis cancer antigen.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Gарв
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           HLA; human leucocyte antigen; melanoma; lung cancer; head cancer;
neck cancer; MAGE-3; human.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Indela
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              100.0%; Score 51; DB 8; I 100.0%; Pred. No. 1.8e+06; iive 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Human HLA-B18 binding MAGE-3 peptide #11.
                                                                                                                                       Disclosure, SEQ ID NO 29; 137pp; English
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Claim 22; SEQ ID NO 12; 15pp; English
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADI19869 standard; peptide; 9 AA.
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(PANI/) PANICHELLI C.
(BCON/) BCON-FALLEUR T
(BRUG/) BRUGGEN P V D.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             BILSBOROUGH J.
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Best Local Similarity
9; Conserv?
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RESULT 7

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The present invention relates to an isolated peptide which binds to a major histocompatibility complex (MHC) molecule to form a complex that is recognized by a cytolytic T-cell which recognizes and lyses cells presenting complexes of HLA-A2 molecules comprising a sequence of 9 amino acids fully defined in the specification, with the proviso that the peptide is not the peptide of S17 itself. Specifically claimed is an HLA-binding peptide comprising a sequence of 9 amino acids fully defined in the specification. The composition and methods are useful for diagnosing or treating cancer, particularly melanoma. The present sequence
                                                                                                                                                                                                                                                                                                                                                          New isolated SSX-2 and SSX-2-related peptides that bind to human leukocyte antigen (HLA) molecules, useful for diagnosing or treating cancer, particularly melanoma.
                 major histocompatibility complex; MHC; cytolytic T-cell; HLA-A2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    cytostatic; gene therapy; vaccine; cancer; immune response.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       100.0%; Score 51; DB 8; Length 9; 100.0%; Pred. No. 1.8e+06;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Melanoma cell line MAGE gene encoded HLA-A2 peptide
                                                                                                                                                                                                                                         (LUDW-) LUDWIG INST CANCER RES.
(TORR-) TORREY PINES INST MOLECULAR STUDIES.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                Example 8; SEQ ID NO 34; 20pp; English.
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                                                                                                                                                                                                                                                                                            Pinilla C;
                                                                                                                                                                         23-JUL-2003; 2003WO-US023306.
                                                                                                                                                                                                         31-JUL-2002; 2002US-0400076P.
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                                   Cytostatic, cancer.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Local Similarity
                                                                                                       WO2004011483-A2
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Sequence 9 AA;
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                                                                                                                                       05-FEB-2004
                                                                       Synthetic.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Treating subject with pathological condition having human leukocyte antigen-Cw6 molecules presented on cell surface by administering peptides to subject for generating immunologically active response against cells.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       The present invention relates to peptides which form immunologically active complexes with MHC molecules. The invention is useful for treating as abject with pathological condition such as cancer which is melanoma by presenting HLA-CW6 molecules on the cell surface. The present sequence is human HLA-CW6 binding MAGE-1 peptide.
                                                                                                                                                                                                                                                                                                                            major histocompatibility; therapy; cancer; melanoma; HLA-Cw6; human;
                                     Gaps
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 Length 9;
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100.0%; Pred. No. 1.8e+06;
tive 0; Mismatches 0;
Score 51; DB 8; I
Pred. No. 1.8e+06;
0; Mismatches 0;
                                                                                                                                                                                                                                                                                         Human HLA-Cw6 binding MAGE-1 peptide #11.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Example 7; SEQ ID NO 13; 15pp; English.
                                                                                                                                                                                       ADI19026 standard; peptide; 9 AA
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 100.0%;
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Query Match
Best Local Similarity 100.
Matches 9; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (ZHAN/) ZHANG Y.
(TRAV/) TRAVERSARI C.
(BOON/) BOON-FALLEUR T.
(BRUG/) BRUGGEN P V D.
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                                                                     GLYDGMEHL 9
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                              MAGE-1; cytostatic
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                                                                                                                                                                                                                                                                                                                                                                              Homo sapiens
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Matches

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Length 9; Indels

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antimicrobial or cytostatic activity. In addition, the invention may prove useful for the production of a vaccine or for gene therapy. The composition and methods disclosed are useful for preventing or treating infectious diseases or cancer. The present sequence is that of a peptide which was used in the exemplification of the invention.
                                                                                                                                                                                                                                                                                                                                                                                                 MAGE-A10; human; Tumour Rejection Antigen; TRA; Human Leucocyte Antigen; HLA; Major Histocompatibility Complex; MHC; cytolytic T-lymphocyte; CTL; immune response stimulator; prophylaxis; therapy; diagnosis; tumour; cancer; TWF; tumour necrosis factor; vaccine; cytostatic.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Novel polypeptides expressed in tumor cells useful for treating car have an ability to complex with a major histocompatibility complex molecule and comprises a specific unbroken amino acid sequence.
                                                                                                                    100.0%; Score 51; DB 8; L 100.0%; Pred. No. 1.8e+06; tive 0; Mismatches 0;
                                                                                                                                                                                                                                                                                   AAY71489 standard; peptide; 10 AA.
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                                                                                                                                                                                                                                                                                                                                                                         Human MAGE-A10 decapeptide-1.
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Best Local Similarity
Matches 9; Conserv
                                                                                                                       Query Match
Best Local Similarity
                                                                                                                                                                              1 GLYDGMEHL
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                                                                                          Sequence 9 AA;
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Matches
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                                                                                                                                                  The present invention relates to a method of increasing an immune response against a tumour cell, comprising administering to a subject with a tumour an amount of IFN-beta receptor agonist and tumour associated antigen (TRA). The method is useful for increasing an immune response against a tumour cell. The present sequence is a peptide used in
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              This invention relates to a novel hybrid antigen which comprises at least one antigenic domain of an infectious agent or tumour antigen and a binding domain that non-covalently binds to a heat shock protein. The invention may be useful for the production of compounds with an
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   hybrid antigen; antigenic domain; infectious agent; tumour antigen; binding domain; heat shock protein; antimicrobial; cytostatic; vaccine; gene therapy; infectious disease; cancer.
                                                           Increasing an immune response against a tumor cell comprises administering to a subject with a tumor an amount of IFN-beta receptor agonist and tumor associated antigen (TAA).
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                                                                                                                                                                                                                                                                                                                Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          omprising an antigenic domain and improved loomains, useful for preventing or treating
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                                                                                                                                                                                                                                                                                   100.0%; Score 51; DB 8; Length 9; 100.0%; Pred. No. 1.8e+06; ive 0; Mismatches 0; Indels
                                                                                                                                                 present invention relates to a method of increasing
                                                                                                                                                                                                                                                                                                                0; Mismatches
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                                                                                                                                                                                                              response against a tumour cell. The r
the exemplification of the invention.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        New hybrid antigens comprising an shock protein-binding domains, use infectious diseases or cancer.
                                                                                                                       Example 7; Page 47; 80pp; English
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      (MOJA-) MOJAVE THERAPEUTICS INC
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2003US-0463746P.
2003US-0503417P.
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Matches 9, Conservative
      Kurnick JT,
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                                   WPI; 2004-239114/22
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                                                                                                                                                                                                                                                         Sequence 9 AA;
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18-APR-2003; 2
16-SEP-2003; 2
12-PEB-2004; 2
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Boon

De Plaen B,

Brasseur F,

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                                                                                    The patent discloses MAGE-A10 and MAGE-A8 polypeptide, nonapeptide and decapeptide sequences, that function as tumour rejection antigens (TRA8). These peptides are capable of forming a complex with major histocompatibility complex (MIG) molecule type HLA-A2.1 (Human Leucocyte Antigen), that are recognised by T-lymphocytes and elicit an immune response from cytolytic T-lymphocytes (CTL). They function as an immune response stimulator. Tumour rejection antigens are useful in prophylaxis, therapy and diagnosis of tumours and are effective in controlling or preventing tumour growth. The present peptide sequence is the human MAGE-A10 decapeptide. I, that corresponds to residues 254-253 of the MAGE-A10 protein. This peptide can serve as a tumour rejection antigen (TRA) and in combination with adjuvants, can produce vaccines useful for treating a variety of tumours that express MAGE-A10
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Gape
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ö
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Similarity 100.0%; Score 51; DB 3; Length 10; Similarity 100.0%; Pred. No. 0.0069; 9; Conservative 0; Mismatches 0; Indels
Claim 9; Page 37; 80pp; English.
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ADR69766;

g 8

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The patent discloses MAGE-A10 and MAGE-A8 polypeptide, nonapeptide and decapeptide sequences, that function as tumour rejection antigens (TRAs).
These peptides are capable of forming a complex with major histocompatibility complex (MHC) molecule type HLA-A2.1 (Human Leuccoyte histocompatibility complex (MHC) molecule type HLA-A2.1 (Human Leuccoyte Antigen), that are recognised by T-lymphocytes and elicit an immune response from cytolytic T-lymphocytes (CTL). They function as an immune response from cytolytic T-lymphocytes (CTL). They function as an immune response stimulator. Tumour rejection antigens are useful in prophylaxis, therapy and diagnosis of tumours and are effective in controlling or protein, comprising nonapeptides and decapeptides, that serve as tumour rejection antigens (TRAs). The novel TRAs encoded by MAGE-A10 is identified using melanoma call line (LB1751-WEL), stimulated by autologous CTL clone (447A/5) to produce TNF (tumour necrosis factor).
Expression of MAGE-A10 has been detected in a variety of tumours like melanomas, carcinomas of the head and neck, bladder and prostate,
                                                                                                                  MAGE-Al0; human; Tumour Rejection Antigen; TRA; Human Leucocyte Antigen; HLA; Major Histocompatibility Complex; MHC; cytolytic T-lymphocyte; CTL; immune response stimulator; prophylaxis; therapy; diagnosis; tumour; cancer; TNF; tumour necrosis factor; cytostatic.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   myelomas and lung cancer. The only normal tissue expressing MAGE-Al0 is the testis
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Novel polypeptides expressed in tumor cells useful for treating car have an ability to complex with a major histocompatibility complex molecule and comprises a specific unbroken amino acid sequence.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        100.0%; Score 51; DB 3; Length 369; 100.0%; Pred. No. 0.34;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               De Plaen E, Boon T;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        WPI; 2000-412317/35.
N-PSDB; AAD01311, AAD01312, AAD01313.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAB80297 standard; protein; 369 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Brasseur F,
                                                                                                                                                                                                                                                                                                                                                                                                                                                       (LUDW-) LUDWIG INST CANCER RES.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Claim 1; Fig 7; 80pp; English.
                                                                                                                                                                                                                                                                                                                                                                 99WO-IB002018.
                                                                                                                                                                                                                                                                                                                                                                                                            98GB-00026143.
                                      12-OCT-2000 (first entry)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 24-APR-2001 (first entry)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Local Similarity 100.
                                                                                   Human MAGE-A10 protein.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Huang L, Van Pel A,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              σ
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Sequence 369 AA;
                                                                                                                                                                                                                                                                            WO200032769-A2.
                                                                                                                                                                                                                                       sapiens
                                                                                                                                                                                                                                                                                                                                                                 26-NOV-1999;
                                                                                                                                                                                                                                                                                                                                                                                                            27-NOV-1998;
                                                                                                                                                                                                                                                                                                                         08-JUN-2000
AAY71485;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAB80297;
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                                                                                                                                                                                                                                       Homo
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ID AAB8
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AC AAB6
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   This invention relates to a novel hybrid antigen which comprises at least one antigenic domain of an infectious agent or tumour antigen and a binding domain that non-covalently binds to a heat shock protein. The invention may be useful for the production of compounds with an antimicrobial or cytostatic activity. In addition, the invention may prove useful for the production of a vaccine or for gene therapy. The composition and methods disclosed are useful for preventing or treating infectious diseases or cancer. The present sequence is that of a peptide which was used in the exemplification of the invention.
                                                                                                                                                                                                                                                                                                                                        hybrid antigen; antigenic domain; infectious agent; tumour antigen; binding domain; heat shock protein; antimicrobial; cytostatic; vaccine; gene therapy; infectious disease; cancer.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Fletchner J, Prince-Cohane K, Mehta S, Slusarewicz P, Andjelic S;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         New hybrid antigens comprising an antigenic domain and improved heat shock protein-binding domains, useful for preventing or treating infectious diseases or cancer.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ö
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    100.0%; Score 51; DB 8;
100.0%; Pred. No. 0.015;
iive 0; Mismatches 0
                                                                                                                                                                                                                                                                                                  Novel hybrid antigen-related peptide #1346,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Example 11; Page 44; 56pp; English.
                                                                                                                                                                    ADR69766 standard; peptide; 20 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (MOJA-) MOJAVE THERAPEUTICS INC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  2003US-0462469P.
2003US-0463746P.
2003US-0503417P.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     13-FEB-2004; 2004WO-US004340.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 12-FEB-2004; 2004US-00776521.
                                                                                                                                                                                                                                                          (first entry)
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                                                            0
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     WPI; 2004-625768/60.
                                          GLYDGMEHL
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  GLYDGMEHL
                    GLYDGMEHL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               WO2004071457-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Sequence 20 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  11-APR-2003;
18-APR-2003;
16-SEP-2003;
                                                                                                                                                                                                                                                                                                                                                                                                                                Unidentified
                                                                                                                                                                                                                                                       18-NOV-2004
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           13-FEB-2003;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          26-AUG-2004
                                                                                                                                                                                                                                                                                                                                                                                                                                                       Synthetic.
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Barber B;

ADR69766

AAC ADR6

AAC ADR6

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DT 18-1

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DD B Nove

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DD B Nove

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DD B Nove

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DD B Nove

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Gaps

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Human prostate cancer antigen #25.

AAY71485 standard; protein; 369 AA.

RESULT 13

Matches

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The present invention relates to human secreted prostate cancer antigen coding sequences (AAF1274-AAF72789) and proteins (AAB80273-AAB80221).
The coding sequences and proteins of the present invention are useful for preventing, treating or ameliorating a medical condition; and for the diagnosis and treatment of diseases and disorders. Diseases and disorders can be diagnosed and treated include (auto)immune diseases (e.g. graft versus host disease and rheumacoid arthritis), inflammatory and allergic disorders (e.g. asthma), hyperproliferative disorders (e.g. cancers and laukemias), cardiovascular disorders (e.g. stroke), arterial occlusive disorders (e.g. arteriosclerosis), anglogenesis related disorders (e.g. crinopathy and keloid scars), ocular disorders (e.g. glacoma), neurological disorders (e.g. Alzheimer's, Parkinson's disease, epilepsy and Creutzfeld-Jakob disease) and infections caused by bacteria, fungi, viruses or parasites. They may also be useful for wound healing, epithelial cell proliferation, supporting cell cuture, tissue regeneration, birth control and as a food additive or preservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ö
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           New nucleic acid molecule encoding human secreted prostate cancer antigens, useful for the diagnosis and treatment of disorders such as cancer, leukemia and autoimmune disease.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Gaps
|mmunosuppressive; nootropic; neuroprotective; antiviral; vulnerary;
               anticonvulsant; antibacterial; antifungal; antiparasitic; cardiant; immune disorder; cardiovascular disorder; neurological disease; infection; cancer; cytostatic; antiarthritic; antirheumatic; antiathatic; anticonvulsant; vasotropic; vulnerary; human; secreted protein; prostate cancer antigen.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ö
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human bladder cancer associated protein sequence SEQ ID NO:149.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      4; Length 369;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    100.0%; Score 51; DB 4; Length 36
100.0%; Pred. No. 0.34;
cive 0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Claim 11; Page 399-400; 433pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABR48215 standard; protein; 369 AA
                                                                                                                                                                                                                                                                            99US-0144972P.
99US-0148681P.
99US-0149173P.
99US-0158004P.
                                                                                                                                                                                                                                                                                                                                                        05-APR-2000; 2000US-0194689P.
                                                                                                                                                                                                                                         20-JUL-2000; 2000WO-US019666
                                                                                                                                                                                                                                                                                                                                                                                         (HUMA-) HUMAN GENOME SCI INC (ROSE/) ROSEN C A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Ouery Match
Best Local Similarity 100.00
Best Local Similarity 100.00
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                1 GLYDGMEHL 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       WPI; 2001-138554/14.
N-PSDB; AAF72765.
                                                                                                                                                                                                                                                                                                                                                                                                                                                    Birse C;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Sequence 369 AA;
                                                                                                                                                                   40200107476-A1.
                                                                                                                                                                                                                                                                                              13-AUG-1999;
17-AUG-1999;
06-OCT-1999;
                                                                                                                                  Homo sapiens
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         12-JUN-2003
                                                                                                                                                                                                        01-FEB-2001.
                                                                                                                                                                                                                                                                              21-JUL-1999;
                                                                                                                                                                                                                                                                                                                                                                                                                                                    Rosen CA,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABR48215;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            RESULT 15
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABR48215
1D ABR4
XX
AC ABR4
XX
DT 12-J
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DB Huma
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The present invention describes a method for detecting a bladder cancerassociated transcript in a cell from a patient. The method comprises contacting a blological sample from the patient with a polynucleotide that eslectively hybridises to a sequence that is 80 % identical to a cadence the human bladder cancer-associated proteins given in ABR4814 to AC51059 concert the human bladder cancer-associated proteins given in ABR4814 to ABR48242). Bladder cancer-associated proteins given in ABR4814 to ABR48242). In a call in vaccine production. The method can be used in antisense gene therapy and can very concer-associated transcript in a cell from a patient. The method is caseful in diagnosing or treating bladder cancer and in screening for compounds that modulate bladder cancer, such as hormones or antibodies. The nucleic acid molecules from the present invention may be used in various screening and diagnosic methods, and for gene therapy, vaccine and/or antisense/inhibition applications
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                ø
                                                                                                                                                                                                                                                                                                                                                                                                                                                       patient, comprises contacting a biological sample from the patient with bladder cancer-associated polynucleotide or antibody.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Gарв
                                                                                                                                                                                                                                                                                                                                                                                                                                             Detecting a bladder cancer-associated transcript in a cell from a
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ö
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Similarity 100.0%; Score 51; DB 6; Length 369;
Similarity 100.0%; Pred. No. 0.34;
9; Conservative 0; Mismatches 0; Indels
                 Human; bladder cancer; cytostatic; gene therapy; vaccine.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Search completed: November 15, 2005, 15:40:01
Job time : 7.69048 secs
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Claim 10; Page 282; 307pp; English.
                                                                                                                                                                                                                                                                                                               (EOSB-) EOS BIOTECHNOLOGY INC
                                                                                                                                                                                               03-JUL-2001, 2001US-0302814P.
03-AVG-2001, 2001US-0310099P.
08-NOV-2001, 2001US-0343705P.
13-NOV-2001, 2001US-0350666F.
12-APR-2002, 2002US-0372246P.
                                                                                                                                                              03-JUL-2002; 2002WO-US021338
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254 GLYDGMEHL 262
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                    Aziz N;
                                                                                                                                                                                                                                                                                                                                                                                                        N-PSDB; ACC51029
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Sequence 369 AA;
                                                                                        WO2003003906-A2
                                                      Homo sapiens.
                                                                                                                           16-JAN-2003
                                                                                                                                                                                                                                                                                                                                                    Mack DH,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Loc
Matches
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GenCore version 5.1.6
(c) 1993 - 2005 Compugen Ltd.
              Copyright
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OM protein - protein search, using sw model

November 15, 2005, 15:32:22 ; Search time 1.2619 Seconds (without alignments) 686.225 Million cell updates/sec Run on:

US-09-856-812B-42 51 1 GLYDGMEHL 9

Title: Perfect score: Sequence:

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

283416 seqs, 96216763 residues Searched:

Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 200000000

summaries Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 sv

PIR_79:*
1: pir1:*
2: pir2:*
3: pir3:*
4: pir4:* **Database**:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

		æ				
Result No.		Que Mat	Length	DB	۵	ription
-	51	100.0	369	7	I38659	melanoma antigen M
8	41	80.4	242	~	D40590	transcription init
М	40	78.4	419	~	G72396	6-phosphofructokin
4	39	76.5	347	~	138008	melanoma antigen M
S	39	76.5	389	N	S74343	probable aspartate
9	38	74.5	317	ď	138661	melanoma antigen M
7	36	70.6	237	~	S64718	formin-binding pro
80	36	70.6		N	H70793	hypothetical prote
6	36	70.6	575	N	AE3486	
10	35	ъ.	94	~	S26082	U
11	35	9.89	180	N	S57944	ADP-ribosylation f
12	35	9.89	180	N	A23741	ADP-ribosylation f
13	35	68.6	180	~	JC4949	ADP-ribosylation f
14	35	9.89	255	ď	T35217	hypothetical prote
15	35	68.6		~	D89888	ornithine carbamoy
16	35	9.89		~	D72482	probable antibioti
17	35	68.6	~	N	S50675	pre-mRNA splicing
18	34	66.7	138	~	T49060	hypothetical prote
19	34	66.7	179	N	JH0260	ADP-ribosylation f
20	34	66.7	180	Н	S37599	ADP-ribosylation f
21	34	66.7	181	N	B36167	ADP-ribosylation f
22	34	66.7	181	N	A36367	ADP-ribosylation f
23	34	66.7	182	N	C49993	ADP-ribosylation f
24	34	66.7	183	~	D49993	ADP-ribosylation f
25	34	66.7	205		D84527	probable ADP-ribos
26	34	66.7	241	N	AF0818	conserved hypothet
27	34	66.7	245		F69343	d-fer
28	34	66.7	248	~	G65025	DNA
29	34	66.7	248	0	B85893	DNA

probable DNA repli	sigmaB regulation	benzothiadiazole-i	aspartate aminotra	hypothetical prote	hypothetical prote	cobyrinic acid A,C	tryptophanase VCA0	tryptophanase (EC	_	tryptophanase [1mp	phosphate transpor	prophage pi3 prote	formate dehydrogen	probable coatomer	hypothetical prote
F91048	A89999	T06276	AE2412	T01136	F84235	F90184	B82492	WZEC	E91209	H86055	AF3500	B86798	B83966	T38944	T23395
~	~	~	~	N	~	~	~	Н	~	N	~	N	~	~	~
248	333	374	388	405	433	434	472	476	476	476	496	595	782	905	961
66.7	66.7	66.7	66.7	66.7	66.7	66.7	66.7	66.7	66.7	66.7	66.7	66.7	66.7	66.7	66.7
34	34	34	34	34	34	34	34	34	34	34	34	34	34	34	34
30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45

ALIGNMENTS

138659
melanoma antigen MAGE-10 - human
C;Species: Homo sapiens (man)
C;Date: 07-Jun-1996 #sequence_revision 07-Jun-1996 #text_change 09-Jul-2004
C;Accession: 138659
C;Accession: L38659
C; Traversari, C.; Gaforio, J.J.; Szikora, J.P.; De Smet, C.; Břte oon, T.

Immunogenetics 40, 360-369, 1994
A,Title: Structure, chromosomal localization, and expression of 12 genes of the MAGE family.
A,Foceseion: I38659; MUID:95012457; PMID:7927540
A,Accession: I38659
A,Status: preliminary; translated from GB/EMBL/DDBJ
A,Molecule type: DNA
A,Residues: 1-369 <-RES>
A,Cross-references: UNIPROT:P43363; EMBL:U10685; NID:g533510; PIDN:AAA68869.1; PID:g53353
C,Genetics: 1-369 <-RES
A,Gross-references: GDB:331126
A,Gene: GDB:MAGEA10; MAGEIO
A,Cross-references: GDB:331126
A,Hap position: Xq28-Xq28
A,Introns: #status absent
C;Superfamily: tumor associated protein MAGE

Gaps ö Match 100.0%; Score 51; DB 2; Length 369; Local Similarity 100.0%; Pred. No. 0.024; les 9; Conservative 0; Mismatches 0; Indels Query Match Best Local S Matches 9

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1 GLYDGMEHL 9 ठ

254 GLYDGMEHL 262 g

RESULT 2

transcription initiation factor sigma, flagellar-specific - Vibrio parahaemolyticus C,Species: Vibrio parahaemolyticus C,Date: 12-May-1994 #sequence_revision 12-May-1994 #text_change 09-Jul-2004

C)Accession: D40590

R;McCarter, L.L.; Wright, M.B.
J. Bacteriol. 175, 3361-337; 1993

A;Title: Identification of genes encoding components of the swarmer cell flagellar motor A;Title: Identification of genes encoding components of the swarmer cell flagellar motor A;Title: Identification of genes encoding components of the swarmer cell flagellar motor A;Title: Identification of genes encoding components of the swarmer cell flagellar motor A;Title: Identification of genes encoding components of genes encoder average and average and average avera

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A,Title: Evidence for lateral gene transfer between Archaea and Bacteria from genome seq
A,Reference number: A72200, MUID:99287316; PMID:10360571
A,Accesslan: G72396
A,Status: preliminary
A,Molecule type: DNA
A,Residus: 1-419 <ARN>
A,Residus: 1-419 <ARN>
A,Residus: 1-419 <ARN>
A,Residus: 1-410 <ARN
A,Re
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C;Superfamily: pyrophosphate-dependent phosphofructokinase, TM0289 type; 6-phosphofructd
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     A;Cross-references: UNIPROT:Q96TG1; EMBL:X82539; NID:g608992; PIDN:CAA57889.1; PID:g6089
C;Genetics:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           6-phosphofructokinase, pyrophosphate-dependent - Thermotoga maritima (strain MSB8)
C; Species: Thermotoga maritima
C; Species: 11-Jun-1999 #sequence_revision 11-Jun-1999 #text_change 09-Jul-2004
C; Accession: G729; Rsequence_revision 11-Jun-1999 #text_change 09-Jul-2004
R; Nelson, K.E.; Clayton, R.A.; Gill, S.R.; Gwinn, M.L.; Dodson, R.J.; Haft, D.H.; Hic Garrett, M.M.; Stewart, A.M.; Cotton, M.D.; Pratt, M.S.; Phillips, C.A.; Richardson,
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MiAlternate names: MAGE-B1 - human
NiAlternate names: MAGE-B2 protein; MAGE-like protein 1
C;Species: Homo sapiens (man)
C;Date: 01-Mar-1996 #sequence_revision 01-Mar-1996 #text_change 09-Jul-2004
C;Date: 01-Mar-1996 #sequence_revision 01-Mar-1996 #text_change 09-Jul-2004
C;Date: 01-Mar-1996 #sequence_revision 01-Mar-1996 #text_change 09-Jul-2004
R;Muscatelli, F: Walker, A.P.; De Plaen, E.; Stafford, A.N.; Monaco, A.P.
Proc. Natl. Acad. Sci. U.S.A. 92, 4987-4991, 1995
A;Title: Isolation and characterization of a MAGE gene family in the Xp21.3 region.
A;Reference number: I38008; MUID:95281581; PMID:7761436
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Score 41; DB 2; Length 242; Pred. No. 1.5; 0; Mismatches 1; Indels
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A,Cross-references: GDB:635712; OMIM:600619
Map position: Xp21.3-Xp21.3
C,Superfamily: tumor associated protein MAGE
Query Match
Best Local Similarity 87.5%;
Matches 7; Conservative
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Best Local Similarity 77.0.
7, Conservative
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37 GIYDGPKHL 45
                                                                                                                                                                                                                                                                         172 GOYDGMEH 179
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Matches 6, Conserv
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A;Molecule type: mRNA
A;Residues: 1-347 <RES>
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S74343
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probable aspartate transaminase (BC 2.6.1.1) aspC sll0402 [similarity] - Synechocystis sINAlternate names: hypothetical protein sll0402 C;Species: Synechocystis sp. A;Variety: PCC 6803
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   A;Cross-references: UNIPROT:Q55128; EMBL:D64001; GB:AB001339; NID:g1001102; PIDN:BAA10263|
A;Note: the nucleotide seguence was submitted to the EMBL Data Library, June 1996
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C;Date: 0'7-Jun-1996 #sequence revision 07-Jun-1996 #text_change 09-Jul-2004
C;Accession: I38661; I38662; FH1297; PH1298; JC2359; G01446
R;De Plaen, E.; Arden, K.; Traversari, C.; Gaforio, J.J.; Szikora, J.P.; De Smet, C.; Brw
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A;Title: Structure, chromosomal localization, and expression of 12 genes of the MAGE fam1
A;Reference number: I38659; MUID:95012457; PMID:7927540
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.,Experimental source: antigen MAGE-4a
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A; Cross-references: EMBL:U10688; NID:9533516; PIDN:AAA68872.1; PID:9533517
A; Cross-references: EMBL:U10688; NID:9533516; PIDN:AAA68872.1; PID:9533517
A; Cross-references: EMBL:U10688; NID:9533516; PIDN:AAA68872.1; PID:9533517
B; Cross-references: C: van der Bruggen, P:; Luescher, I.F.; Lurquin, C.; Chomez, P.; Van Pel, J. Exp. Med. 176, 1453-1457, 1992
J; Exp. Med. 176, 1453-1457, 1992
A; Attlele: A nonapeptide encoded by human gene MAGE-1 is recognized on HLA-A1 by cytolytic A; Accession: PH1294; MUID:93018875; PMID:1402688
                                                                                                                                                                                                                                                                                   R;Kaneko, T.; Sato, S.; Kotani, H.; Tanaka, A.; Asamizu, B.; Nakamura, Y.; Miyajima, N.; o, K.; Okumura, S.; Shimpo, S.; Takeuchi, C.; Wada, T.; Watanabe, A.; Yamada, M.; Yasuda, DNA Res. 3, 109-136, 1996
A;Title: Sequence analysis of the genome of the unicellular cyanobacterium Synechocystis
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C;Accession: S74343
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C;Keywords: aminotransferase; phosphoprotein; pyridoxal phosphate
F;238/Binding site: pyridoxal phosphate (Lys) (covalent) #status F
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          A;Reference number: S74322; MUID:97061201; PMID:8905231
A;Accession: S74343
A;Status: nucleic acid sequence not shown; translation not shown A;Reacule type: DNA
A;Molecule type: DNA
A;Reacidues: 1-389 <KAN>
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N;Alternate names: MAGE 41 protein; melanoma antigen MAGE-X2
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R; Ding, M.; Beck, R.J.; Keller, C.J.; Fenton, R.G.
Biochem. Biophys. Res. Commun. 202, 549-555, 1994
A; Title: Cloning and analysis of MAGE-1-related genes.
A; Reference number: JC2358; MUID:94311935; PMID:8037761
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A;Molecule type: DNA
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Local Similarity 87.5%; Pred. No. 6.5;
108 7; Conservative 0; Mismatches
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A;Residues: 169-172,'T',174-177 <TRA2>
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A;Residues: 169-177 <TRA1>
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A,Status: preliminary
A,Molecule type: DNA
A,Residues: 1-575 <KUR>
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                                       C, Genetics:
A; Gene: Rv3703c
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C;Species: Mycobacterium tuberculosis
C;Date: 17-010-1998 #sequence_revision 17-Jul-1998 #text_change 09-Jul-2004
R;Dates, R.; Davis, R.; Parkhill, J.; Garnier, T.; Churcher, C.; Harris, D.; Gordon, S;
Connor, R.; Davies, R.; Devlin, K.; Feltwell, T.; Gentles, S.; Hamlin, N.; Holroyd, S.;
Rajandream, M.A.; Rogers, J.; Rutter, S.; Seeger, K.; Whitehead, S.; Barrell, B.G.
A;Authors: Sqares, R.; Sulston, J.E.; Taylor, K.; Whitehead, S.; Barrell, B.G.
A;Accession: H7073
A;Accession: H7073
A;Accession: H7073
A;Accession: H7073
A;Accession: Laber DNA
A;Residues: 1-425 <COL>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   A; Molecule type: mRNA
A; Residues: 1-237 <CHA>
A; Cross-creaces: UNIPROT: Q61053; EMBL: U40751; NID: g1255032; PIDN: AAC52479.1; PID: g125
R; Chan, D.C.; Bedford, M.T.; Leder, P.
B; Chan, D.C.; Bedford, M.T.; Leder, P.
A; Chan, D.C.; Powin binding proteins bear WWP/WW domains that bind proline-rich peptides and A; Reference number: S64711; WUID: 96183189; PMID: 8605874
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        A;Cross-references: UNIPROT:069671; GB:AL022121; GB:AL123456; NID:93261559; PIDN:CAA1802
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C;Species: Mus musculus (house mouse)
C;Species: Mus musculus (house mouse)
C;Accession: S64718; S64711
R;Chan, D.C.; Bedford, M.T.; Leder, P.
submitted to the EMBL Data Library, November 1995
A;Reference number: S64718
A;Accession: S64718
A;Accession: JC2359
A;Rolecule type: mXNA
A;Residues: 1-172.'',174-306,'Q',308-317 <DIN>
A;Residues: 1-172.''',174-306,'Q',308-317 <DIN>
A;Residues: 1-172.''',174-306,'Q',308-317 <DIN>
A;Residues: 1-172.''',174-306,'Q',308-317 <DINSAA19007.1; PID:g499124
A;Cross-references: melanoma cell line DMISO
C;Genetics: melanoma cell line DMISO
C;Genetics: GDB:MAGEA4; MAGEA.'' MAGE-X2
A;Gene: GDB:MAGEA4; MAGEA.'' MAGE-X2
A;Gene: GDB:MAGEA4; MAGEA28
A;Map position: XG28-XG38
A;Map position: XG28-XG38
A;Introns: #etatus absent
C;Superfamily: tumor associated protein MAGE
C;Superfamily: tumor associated protein predicted
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                                                                                                                                                                                                                                                                                                                                                                                                                         Length 317;
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A;Residues: 170-191,'S',193-205,'P',207-216,'V',218-222 <CHW>
A;Cross-references: EMBL:U40751
                                                                                                                                                                                                                                                                                                                                                                                                                      h Similarity 75.0%; Pred. No. 8.1; 6; Conservative 1; Mismatches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity 75.0
Matches 6; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    230 GVYDGREH 237
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Best Local Similarity
Matches 6; Conserv
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Ribelvecchio, V.G.; Kapatral, V.; Redkar, R.J.; Patra, G.; Mujer, C.; Los, T.; Ivanova, P.; Malvecchio, V.G.; Kapatral, V.; Redkar, R.J.; P.H.; Hagius, S.; O'Callaghan, D.; Letessk Proc. Natl. Acad. Sci. U.S.A. 99, 443-448, 2002
Affilte: The genome sequence of the facultative intracellular pathogen Brucella melitens A; Reference number: AD3252; PMID:11756688
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          A, Cross-references: UNIPROT: Q8YEK3; GB: AE008917; PIDN: AAL53056.1; PID: g17983917; GSPDB: CA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ribosomal protein S19 - Euglena gracilis chloroplast
C;Species: chloroplast Euglena gracilis
C;Species: chloroplast Euglena gracilis
C;Date: 3.1-Dec-1993 #sequence.revieton 31-Dec-1993 #text_change 09-Jul-2004
C;Accession: S26082; S34522; S34889
R;Christopher, D.A.; Cushman, J.C.; Price, C.A.; Hallick, R.B.
Curr. Genet. 14, 275-286, 1988
A;Title: Organization of ribosomal protein genes rpl23, rpl2, rps19, rpl22 and rps3 on that Reference number: S26080; MUID:89063445; PMID:3143485
A;Accession: S26082
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Meabduse: 1-94 cCHR.
A; Reabduse: 1-94 cCHR.
A; Cross-references: UNIRROT: P19170; EMBL: Z11874; NID: 914353; PIDN: CAA77918.1; PID: 914368
A; Cross-references: UNIRROT: P1910; EMBL: Z11874; NID: 914353; PIDN: CAA77918.1; PID: 914368
R; Hallick, R.B.; Hong, L.; Drager, R.G.; Favreau, M.; Monfort, A.; Orsat, B.; Splelmann, Bubmitted to the EMBL Data Library, January 1993
A; Description: The complete sequence of the Euglena gracilis chloroplast genome (tentatham A; Reference number: S34494
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A; Residues: 1-94 < HALI>
A; Creatdues: 1-94 < HALI>
A; Creatdues: 1-94 < HALI>
A; Creatdues: 1-94 < HALIS
A; Creatdues: 1-94 < HALIS
B; Hallick, R.B.; Hong, L.; Drager, R.G.; Ravreau, M.R.; Monfort, A.; Oreat, B.; Spielman.
Nucleic Acids Res. 21, 3537-3544, 1993
Nucleic Acids Res. 21, 3537-3544, 1993
A; Title: Complete sequence of Euglena gracilis chloroplast DNA.
A; Reference number: $34862; MUID:93347989; PMID:8346031
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     hypothetical cytosolic protein BMEI1875 [imported] - Brucella melitensis (strain 16M) C;Species: Brucella melitensis
C;Species: Brucella melitensis
C;Date: 01-Feb-2002 #sequence_revision 01-Feb-2002 #text_change 09-Jul-2004
C;Accession: AE3486
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                                                                                                                                                    Length 425
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                                                                                                                                                                                                                        2; Indels
                                                                                                                                            Score 36; DB 2;
Pred. No. 29;
0; Mismatches
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Pred. No. 41;
0; Mismatches
A, Experimental source: strain H37Rv
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity 75.0%;
Matches 6; Conservative
                                                                                                                                            Query Match
Best Local Similarity 75.0%;
Matches 6; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               558 GLYDGANH 565
                                                                                                                                                                                                                                                                                                                                                                                73 GLYDAFEH 80
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をごうない かいこ

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F;2/Modified site: myristylated amino end (Gly) (in mature form) #status predicted
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                                                                              Query Match
Best Local Similarity 66.7
Matches 6; Conservative
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Matches 6; Conservative
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GLYDGLDWL 173
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GLYDGLDWL 173
                                                                                                                                                                                                                             1 GLYDGMEHL 9
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|S1 GLFDGME 157
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Best Local Similarity
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A;Residues: 1-180 <HOS>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         A; Accession: JC4949
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D89888
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C;Species: Gallue gallus (chicken)
C;Species: Gallue gallus (chicken)
C;Species: 13-Jan-1996 #sequence_revision 01-Mar-1996 #text_change 09-Jul-2004
C;Accession: S57944
R;Konkel, D.A.; Song, S.K.
Rsibmitted to the EMBL Data Library, October 1990
A;Description: Nucleotide sequence of a putative chicken ADP-ribosylation factor cDNA.
A;Reference number: S57944
A;Reference number: S57944
A;Reference number: S57944
A;Residues: preliminary
A;Molecule type: mRNA
A;Residues: 1-180 <KON>
A;Residues: 1-180 <KON>
C;Reywords: blocked amino end; lipoprotein; myristylation; nucleotide binding; P-loop
C;Superfamily: ADP-ribosylation factor
C;Keywords: blocked amino end; lipoprotein; myristylation; nucleotide-binding motif B
F;89-94/Region: nucleotide-binding motif B
F;89-94/Region: nucleotide-binding motif B
F;126-129/Region: qTP-binding NXXD motif
F;2/Modified site: myristylated amino end (Gly) (in mature form) #status predicted
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A23741
ADP-Thosylation factor 5 - human
C;Species: Homo sapiens (man)
C;Date: 30-Dec-1991 #sequence_revision 30-Dec-1991 #text_change 09-Jul-2004
C;Accession: A23741
R;Tsuchiya, M.; Price, S.R.; Tsai, S.C.; Moss, J.; Vaughan, M.
J. Biol. Chem. 266, 2772-2777, 1991
A;Tsuchiya, M.; Price, S.R.; Tsai, S.C.; Moss, J.; Vaughan, M.
A;Tsuchiya, M.; Price, S.R.; Tsai, S.C.; Moss, J.; Vaughan, M.
A;Tsuchiya, M.; Price, S.R.; Tsai, S.C.; Moss, J.; Vaughan, M.
A;Tsuchiya, M.; Price, S.R.; Tsai, S.C.; Moss, J.; Vaughan, M.
A;Tsuchiya, M.; Price, S.R.; Tsai, S.C.; Moss, J.; Vaughan, M.
A;Accession: A23741
A;Accession: A23741
A;Accession: A23741
A;Residues: 1-180 CTSU-A37; GB:MS7567; NID:g178986; PIDN:AAA90927.1; PID:g178987
A;Cross-references: GDB:136902; OMIM:103188
A;Gross-references: GDB:136902; OMIM:103188
A;Hap position: 17q12-17q21
C;Superfamily: ADP-ribosylation factor
C;Su
A;Cross-references: EMBL:X70810; NID:g415327; PIDN:CAA50101.1; PID:g415757
A;Note: the nucleotide sequence was submitted to the EMBL Data Library, January 1993
C;Genetis rps19
A;Genetis rps19
A;Genes: chloroplast
A;Genes: chloroplast
C;Superfamily: ribosomal protein S19/S15
C;Keywords: chloroplast; protein biosynthesis; ribosome
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                                                                                                                                                                                                                                                                                                                          Similarity 55.6%; Score 35; DB 2; Length 94; Similarity 55.6%; Pred. No. 7.8; 5; Conservative 3; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   1; Indels
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Best Local Similarity 66.7
Matches 6; Conservative
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50 GVYNGKEHI 58
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                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity
Matches 5; Conserv
                                                                                                                                                                                                                                                                                                                                     Query Match
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C;Accession: JC4949
R;Hosaka, M.; Toda, K.; Takateu, H.; Torii, S.; Murakami, K.; Nakayama, K.
B;Hosaka, M.; Toda, K.; Takateu, H.; Torii, S.; Murakami, K.; Nakayama, K.
Biochem. 120, 813-819, 1996
A;Hitle: Structure and intracellular localization of mouse ADP-ribosylation factors type
A;Reference number: JC4945; MUID:97103475; PMID:8947846
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Across-references: UNIPROT: P26437; DDBJ: DBJ: DBJ: DBJ: S1265214; PIDN: BAA13494.1; PID: 9156
A; Experimental source: Drain
C; Comment: This protein belongs to class II of ADP-ribosylation factors which are a fami;
C; Comment: This protein belongs to class II of ADP-ribosylation factors which are a fami;
C; Superfamily: ADP-ribosylation factor
C; Keywords: blocked amino end; lipoprotein; myristylation; nucleotide binding motif A (P-loop)
F; 24-31/Region: nucleotide-binding motif B F; 24-31/Region: GPD-binding motif B F; 24-31/Region: GPD-binding NKXD motif B F; 24-31/Region: GPD-binding NKXD motif B F; 24-31/Region: Myristylated amino end (Gly) (in mature form) #status predicted
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    A;Status: preliminary; translated from GB/EMBL/DDBJ
A;Molecule type: DNA
A;Mosidues: 1-25 < SER-
A;Kosidues: 1-25 < SER-
A;Cross-references: UNIRROT:086707; EMBL:AL031515; PIDN:CAA20622.1; GSPDB:GN00070; SCOED!
A;Experimental source: strain A3(2)
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                                                                                                                                                                                                                                                                                                                                                                                                                                          ADP-ribosylation factor 5 - mouse
C;Species: Mus musculus (house mouse)
C;Date: 31-Dec-1996 #sequence_revision 31-Dec-1996 #text_change 09-Jul-2004
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        C;Species: Streptomyces coelicolor
C;Date: 05-Nov-1999 #sequence_revision 05-Nov-1999 #text_change 09-Jul-2004
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Gaps
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A;Reference number: Z21572
A;Accession: T35217
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       68.6%; Score 35; DB 2; Length 255;
85.7%; Pred. No. 25;
tive 1; Mismatches 0; Indels
2; Length 180;
                                                                                  1; Indels
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C;Superfamily: conserved hypothetical protein MJ1163
    68.6%; Score 35; DB 66.7%; Pred. No. 17;
                                                                              2; Mismatches
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ornithine carbamoyltransferase [imported] - Staphylococcus aureus (strain N315)
C;Species: Staphylococcus aureus
C;Date: 10-May-2001 #sequence_revision 10-May-2001 #text_change 09-Jul-2004
C;Accession: D89888
C;Accession: D89888
N; Mizutani-Ui, Y:; Kobayashi, N.; Sawano, T.; Yuzawa, H.; Kobayashi, I.; Cui, L.; Oguc
Ma, A.; Mizutani-Ui, Y:; Kobayashi, N.; Sawano, T.; Inoue, R.; Kato, C.; Sekimizu, K.;
C; Shiba, T.; Hattori, M.; Ogasawara, N.; Hayashi, H.; Hiramatsu, K.
Lancet 357, 1225-1240, 2001
A;Title: Whole genome sequencing of meticillin-resistant Stapylococcus aureus.
A;Reference number: A89758; MUID:21311952; PMID:11418146
A;Residues: D89888
A;Status: preliminary
A;Molecule type: DNA
A;Residues: 1-333 <KUR>
A;Residues: 1-333 <KUR>
A;Genetics:
A;Genetics:
C;Genetics:
A;Genetics:
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Search completed: November 15, 2005, 15:44:46 Job time : 3.2619 secs

1 GLYDGMEH 8 |:|||:|: 99 GMYDGIEY 106

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GenCore version 5.1.6
(c) 1993 - 2005 Compugen Ltd.
           Copyright
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OM protein - protein search, using sw model

November 15, 2005, 15:31:39 ; Search time 5.2619 Seconds (without alignments) 875.864 Million cell updates/sec Run on:

US-09-856-812B-42 51 1 GLYDGMBHL 9

Title: Perfect score: Sequence:

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

1612378 segs, 512079187 residues Searched:

1612378 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 200000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Database :

UniProt_03:*
1: uniprot_sprot:*
2: uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Description	P43363 homo sapien	2 homo	3 shewa	•	O9bun9 homo sapien		Q9wyc5 thermotoga	homo	homo	homo	homo	canie	Q8dgg0 synechococc	Q55128 synechocyst		symbi	P43358 homo sapien	-	picro	O15481 homo sapien	Q8iz00 homo sapien	Q9kyi5 streptomyce	Q9cnp5 pasteurella	-	Q9hku8 thermoplasm	_	Q7ngq2 gloeobacter	Q9d4d2 mus musculu	Q8r659 fusobacteri	_	Q8sqi6 encephalito
ΩI	MAGA HUMAN	096L <u>Z</u> 2	QBECUB	LAFS VIBPA	9BUN9	Q97AP7	Q9WYC5	MGB1 HUMAN	Q96CW8	Q96TG1	Q6FHJ0	Q9TTY4	Qanggo	AAT SYNY3	Q94 <u>E</u> G1	Q67SK0	MAG4 HUMAN	Q147 <u>9</u> 8	Q6L1Z9	MGB4 HUMAN	081200	Q9KY15	RNFE PASMU	MGB2 HUMAN	Q9HK <u>U</u> 8	Q8S9Y6	Q7NGQ2	Q9D4D2	QBR659	Q9ACL1	AM11_ENCCU
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* Query Match	100.0	84.3	82.4	80.4	80.4	78.4	. 78.4	76.5	76.5	76.5	76.5		76.5	76.5	76.5	74.5	74.5	74.5	74.5	74.5	74.5	72.5	72.5	72.5	72.5	72.5	72.5	72.5	٠		
Score	51	43	42	41	41	40	40	39	39	39	39	39	39	39	39	38	38	38	38	38	38	37	37	37	37	37	37	37	37	37	37
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Q88rg3 encephalito P47715 mycoplasma Q6007 bartonella Q61053 mua musculu Q88rw0 encephalito Q67m8 symbiobacte Q7d313 mycobacteri Q6571 mycobacteri Q7tw2 mycobacteri Q8gpn8 streptomyce Q7nts photorhabdu Q6fal7 acinetobact Q8g380 brucella su Q8g380 brucella su
AM12_ENCCU RPOB_MYCGA 06G0E7 06G0E7 081053 081053 08108 07713 081718 07718 07718 07718 0818 0818 0818
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## ALIGNMENTS

RESULT MAGA H ID M	JT 1 HUMAN MAGA HUMAN P43363;	STANDARD;	RD,	PRT,	369 AA.	AA.
222	01-NOV-1995 (Rel. 32, Created) 01-NOV-1995 (Rel. 32, Last sequence update) 05-JUL-2004 (Rel. 44, Last annotation update)	(Rel. 32, (Rel. 32, (Rel. 44,	Created) Last seq Last ann	<pre>:ed) Bequence update) annotation update) . 10 (MACE-10 anticol</pre>	pdat upd	(e) (atte) (atte)
8 8 8	Name=MAGEA10;	Synonymi (Human)	Synonyme=MAGE10; Human)	45441	5 -	. (115811)
888	Bukaryota, Metazoa, Chordata,	etazoa; C	hordata;	Craniat	V 2 2 4	Craniata; Vertebrata; Euteleostomi;
388	NCBI_TaxID=9606;	606;	i sagent t	כמרמזדו	,	
2	SEQUENCE FRO	M N.A.				
¥ :	MEDLINE=9501	2457; Publ	Med=79275	40,	ć	OF THE STANDS
<b>5</b>	de Smet C.,	Brasseur	F., van d	er Bruc	gen	de Smet C., Brasseur F., van der Bruggen P., Lethe B., Lurquin C.,
¥ £	"Structure,	Chomez F	., de bac al locali	zation	and	renee w., boom 1.; l expression of 12 genes of
RI	the MAGE family."; Imminogenetics 40:360-369(1994)	ily.";	-369(1994	_		
Z Z	[2]	20.05	-			
RP	SEQUENCE FROM N.A.	M N.A.				
۲ ۲	TISSUE=Skin;	1	1000			. 1023 (2000) 2426 2520
\$ £	Straugherg R	L. Fein	medeiza//	Grous	Je I.	MEDLINE=2238623/; Fulled=124//332; DOI=10/3/pmg8.24203039; Stramsherg R.L., Peingold B.A., Grouge L.H., Derge J.G.,
æ	Klausner R.D	Collin	F 1. S	agner		Klausner R.D., Collins F.S., Wagner L., Shenmen C.M., Schuler G.D.,
<b>8</b> 8	Altechul S.F	., Zeeber	Bue Bue	TOW K.	. ×	Schaefer C.F., Bhat N.K., I. Wang J., Haich F.
5 2	Diatchenko L	, Marusi	na K., Fa	rmer A	, A	Rubin G.M., Hong L.,
Z:	Stapleton M.	Soares	M.B., Bor	aldo M	D ₄	Casavant T.L., Scheetz T.E.,
<b>§</b> 2	Raha S.S., L	ocuellano	N.A., PE	ceniyu tera G		Abramson R.D., Mullahy S.J.,
Æ	Bosak S.A.,	McEwan P.	J., McKer	nan K.	7	Malek J.A., Gunaratne P.H.,
<b>§</b> §	Kichards S., Villalon D.K	WOTLEY K	D.M., SOC	ergren	B.J.	A A.M., GRY L.J., Huryk S.W.,
S.	Fahey J., He	lton B.,	Ketteman	M., Ma	lan A	A., Rodrigues S., Sanchez A.,
8 8	Whiting M., Rlakeslev R.	Madan A., W., Touch	Young A.	Green	Syche B.D.	enko Y., Bourrard G.G., . Dickbon M.C.,
2	Rodriguez A.	C., Grimw	ood J., S	chmutz		A Rodriguez A.C., Grimwood J., Schmutz J., Myers R.M.,
\$ £	Schnerch A.	X.S.N., K.	rzywinski .E Jone	M.1.,	ָרָאָמָ מאָמָר	laka U., Smallus D.E., farra M.A.,
F.	"Generation	and initi	al analye	is of	nore	than 15,000 full-length human
R.	and mouse cDNA sequences.";	NA sequen	ces.";		,	(0000)
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ខ	developm	ent and t	umor trar	Bforma	ion	development and tumor transformation or aspects of tumor
ខ្ល		ion.	, D	4	Ē	progression.
ខ	such as	melanoma,	head and	neck i	aguan	ilosob statititi bapiessed in many tumbis of several type, such as melanoma, head and neck squamous cell carcinoma, lung
ខ្ល	carcinom	a and bre	ast carci	noma, l	out n	not in normal tissues except
ខូខូ	for test -1- SIMILARI	for testes and placenta. SIMILARITY: Contains 1 MAGE domain.	acenta. ins 1 MAC	E doma:	Ë.	
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Tats Bearleage 14 (1028) DOI=10.1038/ng1285; C. Tiss Bearleage 14 (10202) DOI=10.1038/ng1285; C. Tiss Bearleau A., Hayashi K., Sato H., Nagai K., Kimura K., Makita H., Sekine M., Obayashi M., Nishi T., Shibahara T., Tanaka T., Ishii S., Sakine M., Obayashi M., Nishi T., Shibahara T., Tanaka T., Ishii S., Amamoto J., Saito K., Kawai Y., Isono Y., Nakamura Y., Nagahari K., Murakami K., Yasuda T., Iwayanagi T., Wagatsuma M., Shiratori A., Saudo H., Hosoiri T., Kautu Y., Kondaira H., Kondo H., Sugawara M., Saudo H., Hosoiri T., Yamashira H., Kinkawa E., Omura Y., Abe K., Ishibashi T., Yamashira H., Murakawa K., Fulimori K., Anai H., Kimata M., Watanabe S., Yosida M., Hotuta T., Kusano J., Anai H., Kimata M., Watanabe S., Yosida M., Hotuta T., Kusano J., Ranehori K., Takahashi-Fujii A., Hara H., Tanase T., Nomura Y., Anehori K., Yuuki H., Oshima A., Sasaki N., Anita M., Imose N., Musashino K., Yuuki H., Oshima A., Sasaki N., Anita M., Sano S., Noshikawa Y., Matanabe T., Sugiyama A., Takemoto M., Kawakami B., Yoshikawa Y., Matanabe T., Sugiyama A., Takemoto M., Kawakami Y., Anamazaki M., Watanabe T., Sugiyama A., Takemoto M., Kawakami Y., Amazaki M., Watanabe T., Sugiyama A., Takemoto M., Kawakami Y., Amazaki M., Watanabe T., Sugiyama A., Takemoto M., Chimori Y., Komiyama M., Tashito H., Itena Y., Okamoto S., Kawabera A., Hikiji T., Kobatake N., Inagaki H., Ikema Y., Okamoto S., Rawabate A., Hikiji T., Kobatake N., Inagaki H., Ikema Y., Okamoto S., Rawabate A., Hikiji T., Robatake N., Inagaki H., Matanabe M., Matsumura K., Sanaki M., Sasaki M., Rawakami T., Oyama M., Hata H., Matanabe M., Komatsu T.,
                       This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the Buropean Bioinformatics Institute. There are no restrictions on its mon-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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01-DEC-2001 (TrEMBLrel. 19, Last sequence update)
01-DRC-2003 (TrEMBLrel. 23, Last annotation update)
01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
Hypothetical protein FLJ32965.
Hypothetical protein FLJ32965.
Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostoml;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             100.0%; Score 51; DB 1; Length 369; 100.0%; Pred. No. 0.44;
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SEQUENCE 369 AA; 40766 MW; 16FA3331CAB716A6 CRC64;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Antigen, Multigene family, Tumor antigen. DOMAIN 134 333 MAGE.
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Genew; HGNC:6797; MAGEA10.
H-InvDB; HIX0017116; -.
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InterPro; IPR002190; MAGE.
Pfam; PF01454; MAGE; 1.
PROSITE; PS50838; MAGE; 1.
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Matches 9; Conservative
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096LZ2;
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Mizushima-Sugano J., Satoh T., Shirai Y., Takahashi Y., Nakagawa K., Okumura K., Nagase T., Nomura N., Kikuchi H., Masuho Y., Yamashita R., Nakai K., Yada T., Nakamura Y., Ohara O., Isogai T., Sugano S.; "Complete sequencing and characterization of 21,243 full-length human
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Heidelberg J.F., Paulsen I.T., Nelson K.E., Gaidos E.J., Nelson W.C., Read T.D., Eisen J.A., Seshadri R., Ward N.L., Methe B.A., Clayton R.A., Meyer T., Tsapin A., Scott J., Beanan M.J., Brinkac L.M., Daugherty S.C., DeBoy R.T., Dodson R.J., Durkin A.S., Haft D.H., Kolonay J.F., Madipu R., Peterson J.D., Umayam L.A., White O., Wolf A.M., Vamathevan J.J., Weidman J.F., Impraim M., Uterback T.R., McDonald L.A., Feldblyum T.V., Smith H.O., Venter J.C., Nealson K.H., Ferser C.M.; "Genome sequence of the dissimilatory metal ion-reducing bacterium
                                                                                                                                                                                                                                                                                                                                                                Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Bacteria; Proteobacteria; Gammaproteobacteria; Alteromonadales;
Shewanellaceae; Shewanella.
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                                                                                                                                                                                                                                                                                                        84.3%; Score 43; DB 2; Length 347; 87.5%; Pred. No. 13; 0; Indels ative 1; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           82.4%; Score 42; DB 2; Length 330; 77.8%; Pred. No. 18; tive 1; Mismatches 1; Indels
                                                                                                                            Nat. Genet. 36:40-45(2004).

EMBL, AKO57527; BAB71522.1; -.

InterPro; IPR002190; MAGE.

Pfam; PF01454; MAGE; 1.

PROSITE; PS5038; MAGE; 1.

SEQUENCE 347 AA; 38878 MW; 3B52411D840D873F CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              STRAIN=MR-1;
MEDLINE=22297686; PubMed=12368813; DOI=10.1038/nbt749;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              330 AA; 37462 MW; 9B2F0235AF427242 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           01-MAR-2003 (TrEMBLrel. 23, Created)
01-MAR-2003 (TrEMBLrel. 23, Last sequence update)
01-UNN-2003 (TrEMBLrel. 24, Last annotation update)
Hypothetical protein S03025.
OrderedLocusNames=S03025;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           (Rel. 27, Created)
(Rel. 27, Last sequence update)
(Rel. 44, Last annotation update)
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Nat. Biotechnol. 20:1118-1123 (2002).
RMBL; AE015739; AAN56037.1; -.
TIGR; S03025; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    InterPro; IPR000801; Bsterase put. Pfam; PF00756; Esterase; 1.
                                                                                                                                                                                                                                                                                   Ouery Match
Best Local Similarity 87.5.
7; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                    231 GLYDGIEH 238
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SEQUENCE 330 AA;
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01-OCT-1993
05-JUL-2004
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LAFS VIBPA
ID LAFS VIB:
AC Q03474;
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QBECUB
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MEDLINE=2238825; PubMed=12477932; DOI=10.1073/pnas.242603899;
MEDLINE=22388257; PubMed=12477932; DOI=10.1073/pnas.242603899;
Attaubberg R.D., Cellins F.S., Wagner L., Shemmen C.M., Schuler G.D.,
Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
A blatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
Biatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
Brownstein M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
Brownstein M.J., Usdin T.B., Tooshiyuki S., Carninci P., Prange C.,
Brownstein M.J., Usdin T.B., Tooshiyuki S., Carninci P., Prange C.,
Brownstein M.J., Usdin T.B., Tooshiyuki S., Carninci P., Prange C.,
Brownstein M.J., WcKernan K.J., Malek J.J., Hulyk S.W.,
Brichards S., Worley K.C., Hals, Soarcla A.N., Gabbs R.A.,
A Nilalon D.K., Muzny D.M., Sodergren B.J., Lu X., Gibbs R.A.,
Whiting M., Madan A., Young A.C., Shevchenko Y., Boulfard G.G.,
Whiting M., Touchman J.W., Green B.D., Dickson M.C.,
Rodriguez A.C., Grimwood J., Schmutz J., Myers R.M., Butterfield Y.S.,
Krzywinski M.I., Skalska U., Smailus D.E., Schnerch A., Schein J.E.,
Jones S.J., Marra M.A.,
"Generation and initial analysis of more than 15,000 full-length human and mouse cDNA sequences."
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Kalnine N., Chen X., Rolfs A., Halleck A., Hines L., Elsenstein S.,
Koundinya M., Raphael J., Moreira D., Kelley T., LaBaer J., Lin Y.,
Phelan M., Farmer A.,
                                                                                                                                                                            Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi,
Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       2; Length 318;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Strausberg R.;
Submitted (FEB-2001) to the EMBL/GenBank/DDBJ databases.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Strausberg R.;
Submitted (AUG-2001) to the EMBL/GenBank/DDBJ databases.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Submitted (MAY-2003) to the EMBL/GenBank/DDBJ databases
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   318 AA; 35214 MW; EA02CIFB42F6C080 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       roc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
                                                    Last sequence update)
Last annotation update)
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01-OCT-2001 (TrEMBLrel. 18, Last sequence update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     11arity 87.5%; Pred. No. 27; Conservative 0; Mismatches
                               Created)
                            01-JUN-2001 (TrEMBLrel. 17,
01-JUN-2001 (TrEMBLrel. 17,
25-OCT-2004 (TrEMBLrel. 28,
Melanoma antigen, family A,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 EMBL; BC002455; AAH02455.1;
EMBL; BT007340; AAP36004.1;
EMBL; BC012744; AAH12744.1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             InterPro; IPR002190; MAGE.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Pfam, PF01454; MAGE, 1.
PROSITE, PS50838; MAGE,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          232 GLYDGREH 239
                                                                                                                                                    Homo sapiens (Human)
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les 7; Conserv
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                                                                                                                                                                                                                              WCBI_TaxID=9606;
                                                                                                                                Name=MAGEAB;
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Q97AP7;
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Matches
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ID 099
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the buspean Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                 STRAIN-RIMD 2210633 / Serotype 03:K6;

MEDLINE-22508454; PubMed=12620739; DOI=10.1016/S0140-6736(03)12659-1;

Makino K., Oshima K., Kurokawa K., Yokoyama K., Uda T., Tagomori K.,

Mijima Y., Najima M., Nakano M., Yamashita A., Kibota Y.,

Yagunaga T., Honda T., Shinagawa H., Hattori M., Iida T.,

"Genome sequence of Vibrio parahaemolyticus: a pathogenic mechanism
distinct from that of V. cholerae.";

Lancet 361:743-749(2003).

--- FUNCTION: Sigma factors are initiation factors that promote the
attachment of RNA polymerase to specific initiation sites and are
then released. This alternative sigma factor is specific for the
flagellin gene (filc) expression.

--- INDUCTION: Under conditions in which the polar flagellum is not
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           55 68 Polymerase core binding (Potential).
211 230 H-T-H motif (By similarity).
242 AA; 27835 MW; 39A3C7152DE2FC95 CRC64;
RNA polymerase sigma factor for flagellar operon (Sigma-F factor)
                                                                                                                                                                                                                                                                                               "Identification of genes encoding components of the swarmer cell flagellar motor and propeller and a sigma factor controlling differentiation of Vibrio parabaemolyticus."; U. Bacteriol. 175:3361-3371(1993).
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Sigma factor; Transcription regulation.
                                                                              Vibrio parahaemolyticus.
astoria, Proteobacteria, Gammaproteobacteria, Vibrionales,
Vibrionaceae, Vibrio.
VCBL_TaxID=670;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 80.4%; Score 41; DB 1; Length 242; 87.5%; Pred. No. 21; ive 0; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     functional.
-!- SIMILARITY: Belongs to the sigma-70 factor family.
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InterPro; IPR007627; Sigma70_r2.
InterPro; IPR007630; Sigma70_r4.
InterPro; IPR000943; Sigma 70.
Pfam; PP04542; Sigma70_r2; 1.
Pfam; PP04545; Sigma70_r2; 1.
PRINTS; PR00046; SIGMA70_r4; I.
PROSITE; PS00715; SIGMA70_1; FALSE_NEG.
                                    (Sigma-27).
Name=lafS; OrderedLocusNames=VPA1555;
                                                                                                                                                                                                                                                     MEDLINE=93273702; PubMed=8501040;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          EMBL; US2957; AAB07356.1; -.
EMBL; AP005089; BAC62898.1; -.
PIR; D40590; D40590.
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                                                                                                                                                                                                                                                                                  McCarter L.L., Wright M.E.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match 80.4
Best Local Similarity 87.5
Matches 7; Conservative
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Gapa

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318 AA.

PRT;

PRELIMINARY;

O9BUN9 RESULT 5 Q9BUN9 ID Q9BUN

DNA BIND SEQUENCE DOMAIN

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ProDom; PD000707; Ppfruckinase; 1.
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                                                                                                                                                                                    SEQUENCE FROM N.A.

STATAL=GS21 / DSW 4299 / JCM 9571;

STATAL=GS21 / DSW 4299 / JCM 9571;

MEDLINE=20570466; PubMed=11121031; DOI=10.1073/pnas.97.26.14257;

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Kawashima -Ohya Y., Watanabe K., Yamazaki M., Kanehori K., Kawamoto T.,

Annoshiba T., Yamanoto Y., Aramaki H., Makino K., Suzuki M.;

Annoshiba T., Yamanoto Y., Aramaki H., Makino K., Suzuki M.;

Richaeal adaptation to higher temperatures revealed by genomic sequence of Thermoplasma volcanium.";

Proc. Natl. Acad. Sci. U.S. A. 97:14262(2000).

REMBL; AP000993; BAB59905.1; --.

REMBL; AP000993; BAB59905.1; --.

RO; GO:0016829; F:lyase activity; IEA.

RO; GO:0016829; F:lyase activity; IEA.

RO; GO:0006520; P:amino acid metabolism; IEA.

REMBL; PRO0291; PALP; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               STRAIN-MSB8 / DSM 3109 / ATCC 43589;

KREDLINE-99287316; PubMed=10360571; DOI=10.1038/20601;

ABDLINE-99287316; PubMed=10360571; DOI=10.1038/20601;

ABDLINE-99287316; PubMed=10360571; DOI=10.1038/20601;

ABTATA-MSBOON R.B. C. Garrent M.B., Garrent M.B., Ratt D.H., Hickey E.K., Peresron J.D., Nelson W.C., Ketchum K.A.,

ACCOMAID L.A., Utterback T.R., Malek J.A., Linher K.D., Garrett M.M.,

ACCOMAID L.A., Utterback T.R., Malek J.A., Linher K.D., Garrett M.M.,

ACCOMAID L.A., Sutton G.G., Fleischmann R.D., Eisen J.A., White O.,

Alzberg S.L., Smith H.O., Venter J.C., Fraser C.M.;

The state of Thermotoga maritima.";

BENL; ACCOMAID AAD35377.1; -.

REMBL; ACCOMAID AAD35377.1; -.

REMBL; ACCOMAID AAD35377.1; -.

REMSP: P70826; IKZH.

RIGR; TM0289; LXZH.
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                                                                                          Archaea, Buryarchaeota, Thermoplasmata, Thermoplasmatales,
Thermoplasmataceae, Thermoplasma.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          78.4%; Score 40; DB 2; Length 340; 77.8%; Pred. No. 44;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               2; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Complete proteome; Pyridoxal phosphate.
SEQUENCE 340 AA; 37877 MW; FC2864128896433B CRC64;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        01-NOV-1999 (TrEMBLrel. 12, Created)
01-NOV-1999 (TrEMBLrel. 12, Last sequence update)
01-UDN-2003 (TrEMBLrel. 24, Last annotation update)
6-phosphofructokinase, pyrophosphate-dependent.
OrderedLousnames=TM0289;
01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             0; Mismatches
                    Threonine synthase.
Name=TVG0766521; OrderedLocusNames=TV0763;
Thermoplasma volcanium.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity 77.8
Matches 7; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                PRELIMINARY;
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                                                                                                                                            NCBI_TaxID=50339;
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DDE REPRESENT OF THE PROPERTY 
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This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the European Bioinformatics Institute. There are no restrictions on its mee by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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-I- TISSUE SPRCIFICITY: Expressed only in testis.
                                                                                                                                                                   Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          TISSUE=Testis;
MEDLINE=56081328; PubMed=8535061;
Dabovine B., Zanaria E., Bardoni B., Lisa A., Bordignon C., Russo V.,
Matessi C., Traversari C., Camerino G.;
"A family of rapidly evolving genes from the sex reversal critical
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Muscatelli F., Walker A.P., de Plaen E., Stafford A.N., Monaco A.P.; "Isolation and characterization of a MAGE gene family in the Xp21.3
                                                                                                                                                                                                                                                                                                                                                                                                                                                             MGB1 HUMAN STANDARD; PRT; 347 AA.
P43366; 000601; 075862;
01-NOV-1995 (Rel. 32, Created)
16-CCT-2001 (Rel. 40, Last sequence update)
05-UUL-2004 (Rel. 44, Last annotation update)
05-UUL-2010 (Rel. 44, Last annotation update)
(DSS-AHC critical interval MAGE-B1 antigen) (WAGE-XP antigen)
(DSS-AHC critical interval MAGE superfamily 10) (DAM10).
Name-MAGEB1; Synonyms=MAGEL1, MAGEXP;
Homo saplens (Human).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                ;
                                                                                             Length 419;
                                                                                                                                                                Indels
Complete proteome; Kinase.
SEQUENCE 419 AA; 46464 MW; 4B3FBC75A8410CEC CRC64;
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                                                                                                 ;
7
                                                                                             Score 40; DB
Pred. No. 54;
                                                                                                                                                                2; Mismatches
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EMBL; S80936; AAC97145.1; -.
EMBL; AC005185; AAD10634.1; -.
Genew; HGNC:6808; MAGEB1.
H-INVDB; HXX0016714; -.
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                                                                                                 78.4%;
                                                                                                                                                                   Conservative
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37 GIYDGFKHL 45
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                                                           Query Match
Beet Local Similarity
6, Conserve
                                                                                                                                                                                                                                  1 GLYDGMEHL
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Euteleostomi;

228 GAYDGEEHL 236

InterPro; IPR002190; MAGE

STTTTS

Q96CW8

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SEQUENCE FROM N.A.
MEDLINE-98110575; PubMed-9441743; DOI-10.1006/geno.1997.5052;
Lurquin C., De Smet C., Brasseur F., Muscatelli F., Martelange V.,
De Plaen B., Brasseur R., Monaco A.P., Boon T.,
"Two members of the human MAGES gene family located in Xp21.3 are
expressed in tumors of various histological origins.";
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Mammalia; Butheria; Primates; Catarrhini; Hominidae; Homo.
NCBI_TaxID=9606;
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Halleck A., Ebert L., Mkoundinya M., Schick M., Eisenstein S., Neubert P., Kstrang K., Schatten R., Shen B., Henze S., Mar W., Korn B., Zuo D., Hu Y., LaBaer J.; Shen B., Submitted (JUN-2004) to the EMBL/GenBank/DDBJ databases.

EMBL; CR541762, CAG46562.1;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          76.5%; Score 39; DB 2; Length 347; 77.8%; Pred. No. 70; ive 0; Mismatches 2; Indels
                                                                                                                                                                                                                                                                                                                                          Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Eutele
Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Lurguin C.;
Submitted (JUN-1998) to the EMBL/GenBank/DDBJ databases.
EMBL; U93163; AAC23616.1; -.
PIR; 138008; 138008.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              InterPro; IPR002190; MAGE.
Pfam; PF01454; MAGB; 1.
PROSITE; PS50838; MAGE; 1.
SEQUENCE 347 AA; 39152 MW; 8C54E7ED80C739C0 CRC64;
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PROSITE; PS50838; MAGE; 1.
SEQUENCE 347 AA; 39038 MW; 925E7DB4F19BD25D CRC64;
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Last sequence update)
Last annotation update)
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Last annotation update)
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                                                                                                                   347 AA
                                                                                                                                                                         Created)
                                                                                                                                                                      01-DEC-2001 (TrEMBLrel. 19, 01-DEC-2001 (TrEMBLrel. 19, 01-JUN-2003 (TrEMBLrel. 24,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              05-JUL-2004 (TrEMBLrel. 27, 05-JUL-2004 (TrEMBLrel. 27, 05-JUL-2004 (TrEMBLrel. 27,
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                                                                                                                PRELIMINARY;
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                                                                                                                                                                                                                                                                                         Name=MAGE-B1;
Homo sapiens (Human)
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Best Local Similarity
Matches 7; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                         NCBI_TaxID=9606;
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Riausner R.D., Collins F.S., Wagner L., Shenmen C.M., Schuler G.D.,
Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Hsieh F.,
Diatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
Blatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
Brownstein M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
Brownstein M.J., Usdin T.B., Tooshlyuki S., Carinnoi P., Frange C.,
Brownstein M.J., Usdin T.B., Tooshlyuki S., Carinnoi P., Frange C.,
Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
Bosak S.A., McKwan P.J., McKernan K.J., Makek J.A., Gunstane P.H.,
Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
Villalon D.K., Muzny D.M., Sodergren B.J., Lu X., Gibbs R.A.,
Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
Miking M., Madan A., Young A.C., Schwutz J., Myers R.M., Buterfield Y.S.,
Krzywinski M.I., Skalska U., Smailus D.E., Schmerch A., Schein J.E.,
Generation and initial analysis of more than 15,000 full-length human
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MEDLINE=22388257; PubMed=12477932; DOI=10.1073/pnas.242603899;
                                                                                                                                                                                                                                                  DB 1; Length 347; 70;
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                                                                                                                                                                                                                                                                                                                       2; Indels
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Submitted (SEP-2001) to the EMBL/GenBank/DDBJ databases.
EMBL, BC013772; AAH13772.1; -.
InterPro; IPR002190; MAGE.
Pfam; PF01454; MAGE; 1.
PROSITE; PS50838; MAGE; 1.
PROSITE; PS50838; MAGE; 1.
                                                                                                                   A -> R (in Ref. 1).
G -> S (in Ref. 1).
T -> I (in Ref. 2).
; 538A65E02DC34155 CRC64;
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Last annotation update)
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                                                                                                                                                                                                                                                                                                                    0; Mismatches
                                                                                                                                                                                                                                                                 Score 39;
Pred. No.
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347 AA; 39037 MW;
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77.8%;
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Melanoma antigen, family B,
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Pfam; PF01454; MAGE; 1.
PROSITE; PS50838; MAGE; 1.
Antigen; Multigene family.
DOMAIN 108 307
                                                                                                                                                                                                                                   Query Match
Best Local Similarity 77.5.
T. Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      PRELIMINARY;
                                                                                                             28
172
327
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                                                                                                                                                                                                                                                                                                                                                                              1 GLYDGMEHL 9
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SEQUENCE FROM N.A.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Name=MAGEB1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         01-DEC-2001
01-DEC-2001
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RESULT 9
096CW8
10 096CW8
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RESULT 14

AAT SYNY

12 AAT SYNY

AAT SYNY

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OG 5512

OG 671

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OG 673

OG 6
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Gaps
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                                                                                                                                                                                                                                            delanoma antigen.

Canis familiaris (Dog).

Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Carnivora; Fissipedia; Canidae; Canis.
                                                                                                                                                                                                                                                                                                                                                                                                          STRAIN=breed beagle; TISSUE=Testis;
MEDLINE=20470598; PubMed=11019919;
Ma Z., Khatlani T.S., Ohno K., Sasaki K., Inokuma H., Onishi T.;
"Cloning and sequencing of canine MAGE CDNA.";
Tissue Antigens 56:166-169(2000).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Nakamura Y., Kareko T., Sato S., Ikeuchi M., Katoh H., Sasamoto Watanabe A., Iriguchi M., Kawashima K., Kimura T., Kishida Y., Kiyokawa C., Kohara M., Matsumoto M., Matsumo A., Nakazaki N., Shimpo S., Sugimoto M., Takeuchi C., Yamada M., Tabata S., "Complete genome structure of the thermophilic cyanobacterium Thermosynechococcus elongatus BP-1.";
DNA Res. 9:123-130(2002).
HSSP: QBRR70; 1J32.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 76.5%; Score 39; DB 2; Length 378; 87.5%; Pred. No. 75; ive 0; Mismatches 1; Indels
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Synechococcus elongatus (Thermosynechococcus elongatus)
Bacteria; Cyanobacteria; Chroococcales; Synechococcus.
NCBI_TaxID=32046;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    350838; MAGE; 1.
378 AA; 42214 MW; AD09C6DC8E3C8D57 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  01-MAR-2003 (TrEMBLrel. 23, Created)
1-MAR-2003 (TrEMBLrel. 23, Last sequence update)
01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
Aspartate aminotransferase.
                                                                                                                                                                                                               Last sequence update)
Last annotation update)
                                                                                                                                                  378 AA
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                                                                                                                                                                                             Created)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    EMBL, AF187325; AAF01438.1; -.
InterPro; IPRO02190; MAGE.
Pfam; PF04144; MAGE; 1.
PROSITE; PSS0838; MAGE; 1.
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PRINTS; PR00753; ACCSYNTHASE
                                                                                                                                                                                      01-MAY-2000 (TrEMBLrel. 13, 01-MAY-2000 (TrEMBLrel. 13, 01-MAR-2003 (TrEMBLrel. 23,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Local Similarity 87.5 nes 7; Conservative
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                                                                                                                                                  PRELIMINARY;
                                        228 GAYDGEÉHL 236
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GLYDGMEHL 9
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Matches
                                                                                                       RESULT 12
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This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the Buropean Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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HSSP: ORRATO; 1J32.

InterPro; IPR001176; ACC synthase.

InterPro; IPR004839; Aminotrans I/II.

InterPro; IPR004839; Mainotran I = BS.

Pfam; PF00155; Aminotran I = 1 1.

PRINTS; PR00753; ACCSYNTHASE.

PRINTS; PR00753; ACCSYNTHASE.

PRINTS; PR00759; AA_TRANSFER_CLASS_1; 1.

Aminotransferase; Complete proceeme; Pyridoxal phosphate; Transferase.

238 Pyridoxal phosphate (By similarity).
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-!- SUBUNIT: Homodimer (By similarity).
-!- SUBCELLIVIAR LOCATION: Cytoplasmic (By similarity).
-!- SIMILARITY: Belongs to the class-I pyridoxal-phosphate-dependent aminotransferase family.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Tabata 8.;
"Sequence analysis of the genome of the unicellular cyanobacterium Symechocystis sp. strain PCC6803. II. Sequence determination of the entire genome and assignment of potential protein-coding regions."; DNA Res. 3:109-136(1996).
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MEDLINE=97061201; PubMed=8905231;
Kaneko T., Sato S., Kotani H., Tanaka A., Asamizu E., Nakamura Y.,
Miyajima N., Hirosawa M., Sugiura M., Sasamoto S., Kimura T.,
Hosouchi T., Mateuno A., Muraki A., Nakazaki N., Naruo K., Okumura Shimpo S., Takeuchi C., Wada T., Watanabe A., Yamada M., Yasuda M.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            MEDILINE=96127529; PubMed=8590279; MEDILINE=96127529; PubMed=8590279; MEDILINE=96127529; Tanaka A., Sato S., Kotani H., Sazuka T., Miyajima N., Sugiura M., Tabata S.; Sequence analysis of the genome of the unicellular cyanobacterium Sequence analysis of the genome of the unicellular cyanobacterium sequence analysis of the genome of the genome."; Is Sequence features in the 1 MD region from map positions 64% to 92% of the genome.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         30-MAY-2000 (Rel. 39, Created)
30-MAY-2000 (Rel. 39, Last sequence update)
25-OCT-2004 (Rel. 45, Last annotation update)
Aspartate aminotransferase (EC 2.6.1.1) (Transaminase A) (ASPAT).
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                                                                                                                                     Score 39; DB 2; Length 387;
Pred. No. 77;
0; Mismatches 1; Indels
PROSITE; PS00105; AA TRANSFER CLASS 1; 1. Aminotransferase. Complete proteome; Transferase. SEQUENCE 387 AA; 42123 MW; 6D32D2B740A3CB5F CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Name=aspC; OrderedLocusNames=s110402;
Synechocystis sp. (strain PCC 6803).
Bacteria; Cyanobacteria; Chroococcales; Synechocystis.
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                                                                                                                                                76.5%;
87.5%;
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                                                                                                                                            Query Match
Best Local Similarity
Matches 7; Conserv
                                                                                                                                                                                                                                                                                         2 LYDGMEHL
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Q55128;
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Dubmed=12447438; DoI=10.1038/nature01184; Segurence FROM N.A.
Sagaaki T., Matsumoto K., Sakata K., Baba T., Katayose Y., Sabaki T., Matsumoto T., Sanamoto K., Sakata K., Baba T., Kanamoti H., N. Wi J., Niimura Y., Cheng Z., Nagamura Y., Antonio B.A., Kanamoti H., Hosokawa S., Masukawa M., Arikawa K., Chiden Y., Hayashi M., Okamoto M., Ando T., Aoki H., Arita K., Hamada M., Harda G., Namoto M., Tich S., Itoh T., Itoh Y., Itoh Y., Iwabuchi A., Kamiya K., Kanasawa W., Katagaya W., Nasahima M., Nakamidh Y., Mukai Y., Nakamiti N., Negishi M., Oha I., Oha
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Oryza sativa (japonica cultivar-group).
Eukaryota; Viridiplantea; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
Ehrhartoldeae; Oryzae; Oryza.
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                                                                              76.5%; Score 39; DB 1; Length 389; 87.5%; Pred. No. 78; tive 0; Mismatches 1; Indels
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GO:0009058; P:biosynthesis; IEA.
InterPro; IPR004839; Aminotrans I/II.
InterPro; IPR004838; Mtransf_1_BS.
Pfam; PP00155; Aminotran 1 2; 1.
PROSITE; PS00105; AA TRANSFER CLASS 1; 1.
SEQUENCE 394 AA; 43747 MW; BE843B36Bb6D2762 CRC64;
SQ SEQUENCE 389 AA; 42352 MW; 7706C9C353EA317F CRC64;
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01-DEC-2001 (TrEMBLrel. 19, Last sequence update)
01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
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                                              Query Match
Best Local Similarity 87.5'
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Best Local Similarity 85.79
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             PRELIMINARY;
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C94EG1
ID C94EG1
ID C94EG1
DT C01-DE
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Search completed: November 15, 2005, 15:43:48 Job time : 7.2619 secs

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4268, Ap 3571, Ap 4996, Ap 10, Appli 10, Appli 18, Appli 18, Appli 18, Appli 12, Appli

Sequence Sequence Sequence Sequence

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Run on:

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Query Match
100.0%; Score 51; DB 2; Length 369;
Best Local Similarity 100.0%; Pred. No. 0.092;
Matches 9; Conservative 0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                    Sequence 4, Application US/08773870;
Patent No. 5912143;
GENERAL INFORMATION:
APPLICANT: Bandman, Olga;
APPLICANT: Bandman, Olga;
NUMBER OF SEQUENCES: 6
CORRESPONDENCE ADDRESS:
ADDRESSE: Incyte Pharmaceuticals, Inc.;
STREET: 3174 Porter Drive;
CTY: Palo Alto
STATE: CA
COUNTRY: USA
                                                          US-09-340-812-2
US-08-938-546-4
US-09-340-812-4
US-09-134-001C-4268
US-09-134-001C-4268
US-09-540-236-3571
US-09-936-135-8
US-08-936-135-10
US-08-936-135-18
US-08-936-135-18
                                   JS-09-248-796A-20264
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        COUNTY: USA
ZIF: 94304
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FRATSEQ for Windows Version 2.0
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/773,870
FILING DATE: Herewith
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US/08/773,870
FILING DATE: HOROWATION:
APPLICATION NUMBER: 35
ATTORNEY, AGENT INFORMATION:
NAME: BILLINGS, Lucy J.
REGISTRATION NUMBER: 36,749
REGISTRATION NUMBER: 415-855-0555
                                                                                                                                                                                          US-09-439-711C-10
US-09-439-711C-18
US-08-936-135-12
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     INFORMATION FOR SEQ ID NO: 4:
SEQUENCE CHARACTERISTICS:
LENGTH: 369 amino acids
TYPE: amino acid
STRANDENNESS: single
"nobology: linear
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   inMMEDIATE SOURCE:
LIBRARY: GenBank
CLONE: 533511
US-08-773-870-4
JS-08-773-870-4
 Sequence 29, Appl
Sequence 29, Appl
Sequence 29, Appl
Sequence 29, Appl
Sequence 31, Appli
Sequence 11,71, A
Sequence 11172, A
Sequence 28162, A
Sequence 44, Appl
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Sequence 9310, Ap
Sequence 5, Appli
Sequence 1213, Ap
Sequence 31, Appl
Sequence 31, Appl
Sequence 9891, Ap
Sequence 255, Appl
Sequence 255, Appl
Sequence 2890, Ap
Sequence 2890, Ap
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                                                                                     November 15, 2005, 15:33:16; Search time 1.45238 Seconds (without alignments) 462.579 Million cell updates/sec
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Description
                                                                                                                                                                                                                                                                                                                                                                                       /cgm2 6/ptcdata/1/iaa/5A COMB.pep:*
/cgm2 6/ptcdata/11iaa/5B COMB.pep:*
/cgm2 6/ptcdata/1/iaa/6A COMB.pep:*
/cgm2 6/ptcdata/1/iaa/6B COMB.pep:*
/cgm2 6/ptcdata/1/iaa/PcTUS_COMB.pep:*
/cgm2 6/ptcdata/1/iaa/PcTUS_COMB.pep:*
            GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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US-09-134-001C-2986
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-09-710-279-2890
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                                                                                                                                                                                                                                  513545 seqs, 74649064 residues
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                                                                                                                                                                                                                                                                                                                           Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                              OM protein - protein search, using sw model
                                                                                                                                                                                           BLOSUM62
Gapop 10.0 , Gapext 0.5
                                                                                                                                                                                                                                                                                                                                                                                Issued Patents AA:*
                                                                                                                                                                                                                                                                                     Minimum DB seq length: 0
Maximum DB seq length: 200000000
                                                                                                                                          US-09-856-812B-42
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                                                                                                                                                                   1 GLYDGMEHL 9
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Match Length
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Perfect score:
                                                                                                                                                                                            Scoring table:
                                                                                                                                                                     Sequence:
                                                                                                                                                                                                                                    Searched:
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Result

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Gape

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Sequence 29, Application US/09069226

Patent No. 6013509

GENERAL INFORMATION:
APPLICANT: WARREN, Patrick V.
TITLE OF INVENTION: TRANSAMINASES AND AMINOTRANSFERASES
NUMBER OF SEQUENCES: 32
CORRESPONDENCE 32
CORRESPONDENCE ADDRESS:
ADDRESSEE: CARELLA, BYRNE, BAIN, GILFILLAN,
ADDRESSEE: CACCHI, STEWART & OLSTEIN
STREET: 6 BECKER FARM ROAD
CITY: ROSELAND
STATE: NEW JERSEY
COUNTRY, USA
COUNTRY, USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match 78.4%; Score 40; DB 2; Length 398; Best Local Similarity 75.0%; Pred. No. 12; Matches 6; Conservative 2; Mismatches 0; Indels
COUNTEX: US
ZIP: 92037
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette COMPATION COMPATION COMPATION COMPATION COMPATION DATA:
APPLICATION NUMBER: US/08/646,590B
FILING DATE: 08-May-1996
CLASSIFICATION ATA:
APPLICATION NUMBER: 08/599,171
FILING DATE: 08-FEB-1996
PRIOR APPLICATION NUMBER: PCT/US97/01094
FILING DATE: 21-January-1997
CLASSIFICATION: 435
APPLICATION NUMBER: PCT/US97/01094
FILING DATE: 1-January-1997
CLASSIFICATION NUMBER: 38,347
REFERENCE/DOCKET NUMBER: 38,347
REFERENCE/DOCKET NUMBER: 38,347
TELECHMUNICATION INFORMATION:
TELEPHONE: 619/678-5099
INFORMATION FOR SEQ ID NO: 29:
SEQUENCE CHARACTERISTICS:
LENGTH: 398 amino acide
TYPE: mino acid
TOPLE COMPANIENT NUMBER: TYPE: LENGTH: 398 amino acide
TYPE: mino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    COMPUTER READABLE FORM:
MEDIUM TYPE: 3.5 INCH DISKETTE
COMPUTER: 1BM PS/2
OPERATING SYSTEM: MS-DOS
SOFTWARE: WORD PERFECT 5.1
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/069,226
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/599,171
FILING DATE:
ATTORNEY/AGENT INFORMATION:
NAME: HERRON, CHARLES J.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    MOLECULE TYPE: protein FRAGMENT TYPE: internal
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 :||||||:
210 IYDGMEHV 217
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                                                                                                                                                                                                          GENERAL INFORMATION:
APPLICANT: WARREN, PALTICK V.
TITLE OF INFORMATION:
TITLE OF INFORMATION:
TORRESPONDENCE 32
CORRESPONDENCE ADDRESS:
ADDRESSEE: CARELLA, BYRNE, BAIN, GILFILLAN,
ADDRESSEE: CECCHI, STEWART & OLSTEIN
STREET: 6 BECKER FARM ROAD
CITY: ROSELAND
STATE: MEW JERSEY
COUNTRY: USA
ZIP: 07068
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Fatent No. 5952283
GENERAL INFORMATION:
APPLICANT: Warren, Patrick V.
TITLE OF INVENTION: TRANSAMINASES AND AMINOTRANSFERASES
NUMBER OF SEQUENCES: 42
CORRESPONDENCE ADDRESS:
STREET: ADDRESSE:
STREET: La Jolla
STATE: CA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   78.4%; Score 40; DB 2; Length 398; 75.0%; Pred. No. 12; cive 2; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                            COMPUTER READABLE FORM:
MEDIUM TYPE: 3.5 INCH DISKETTE
COMPUTER: 1BM PS/2
OPERATING SYSTEM: MS-DOS
SOFTWARE: WORD PERFECT 5.1
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/599,171A
FILING DATE: Concurrently
CLASSIPICATION 1435
PRIOR APPLICATION DATA:
APPLICATION NUMBER:
                                                                                                                                                US-08-599-171A-29
; Sequence 29, Application US/08599171A
; Patent No. 5814473
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            PEPLICATION: 435
FILING DATE:
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAWE: HERRON, CHARLES J.
REGISTRATION NUMBER: 28,019
REGISTRATION NUMBER: 3314(
TELEPHONE: 201-994-1700
TELEPRAX: 201-994-1700
TELEFRAX: 201-994-1744
INFORMATION FOR SEQ ID NO: 29:
SEQUENCE CHARACTERISTICS:
LENGTH: 398 AMINO ACIDS
TYPE: AMINO ACID
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity 75.07
Page 6; Conservative
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MOLECULE TYPE: PROTEIN
US-08-599-171A-29
                                                          254 GLYDGMEHL 262
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210 IYDGMEHV 217
                1 GLYDGMEHL 9
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; Sequence 29, Application US/09412184
; Sequence 29, Application US/09412184
; Patent No. 6568188
; Patent No. 6568188
; Patent No. 6568188
; Patent No. 6568188
; APPLICANT: Warren, Patrick V.
; TITLE OF INVENTION: TRANSAMINASES AND AMINOTRANSFERASES
; NUMBER OF SEQUENCES: 42
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Fish & Richardson, P.C.
; STREET: A225 Executive Square, Suite 1400
; CITY: La Jolla
; STATE: CA
                                                                                                                                                                                                                                                                                                                                  Score 40; DB 3; Length 398;
Pred. No. 12;
                                                                                                                                                                                                                                                                                                                                                                                               0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           CITY: La Jolla
STATE: CA
COUNTY: La Jolla
ZIP: 92037
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: Diskette
COMPUTER: Diskette
COMPUTER: DISKETTE
CORFATING SYSTEM: Windows95
SOFTWARE: PASISEO for Windows Version 2.0
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/412,184
FILING DATE:
CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US/08/646,590
FILING DATE: 08-May-1996
APPLICATION NUMBER: 08/599,171
FILING DATE: 09-FEB-1996
APPLICATION NUMBER: PCT/US97/01094
FILING DATE: 21-January-1997
CLASSIFICATION NUMBER: PCT/US97/01094
FILING DATE: 21-January-1997
CLASSIFICATION NUMBER: BCT/US97/01094
FILING DATE: 21-January-1997
ATTONERAL Haile, Ph.D., Lisa A.
REGISTATION NUMBER: BCT/US97/01094
FELECOMMUNICATION INPORMATION:
TELECOMMUNICATION INPORMATION:
TELECOMMUNICATION INPORMATION:
TELECOMMUNICATION INPORMATION:
                                                                                                                                                                                                                                                                                                                                                                                               2; Mismatches
                REFERENCE/DOCKET NUMBER: 331400-38
TELECOMMUNICATION INFORMATION:
TELEPHONE: 201-994-1740
TELEPHONE: 201-994-1744
INFORMATION FOR SEQ ID NO: 29:
SEQUENCE CHARACTERISTICS:
LENGTH: 398 AMINO ACIDS
TYPE: AMINO ACID
TYPE: AMINO ACID
MOLECTLE TYPE: PROTEIN
US-09-069-226-29
REGISTRATION NUMBER: 28,019
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                                                                                                                                                                                                                                                                                                                                       78.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     : 398 amino acids
amino acid
                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity 75.0
Matches 6; Conservative
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FRAGMENT TYPE: internal
US-09-412-184-29
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210 IYDGMEHV 217
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78.4%; Score 40; DB 3; Length 398;

Query Match

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Sequence 22159, Application US/09248796A

Factor No. 6747137

GENERAL INFORMATION:
GENERAL INFORMATION:
TITLE OF INVENTION: FOR DIAGNOSTICS AND THERAPEUTICS
TITLE OF INVENTION: FOR DIAGNOSTICS AND THERAPEUTICS
TITLE OF INVENTION: WHERE 10799-02-12
CURRENT FILING DATE: 1999-02-12
PRIOR APPLICATION NUMBER: US 60/074,725
PRIOR APPLICATION NUMBER: US 60/096,409
FRIOR FILING DATE: 1998-02-13
FRIOR APPLICATION NUMBER: US 60/096,409
FRIOR SPELICATION NUMBER: US 60/096,409
FRIOR APPLICATION NUMBER: US 60/096,409
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Pred. No. 11;
1; Mismatches 1; Indels
                             Indels
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US-08-773-870-3

Sequence 3, Application US/08773870

Patent No. 5912143

GENERAL INFORMATION:
APPLICANT: Bandman, Olga
APPLICANT: Goli, Surya K.

TITLE OF INVENTION: NOVEL HUMAN MAGE-LIKE PROTEIN
NUMBERS OF SEQUENCES: 6

CORRESPONDENCE ADDRESS: 6

CORRESPONDENCE ADDRESS: 6

CITY: Palo Alto
CITY: Palo Alto
CITY: VSA
ZIP.
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COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FastSEQ for Windows Version 2.0
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/773,870
FILING DATE: Herewith
CLASSIFICATION: 435
75.0%; Pred. No. 12; tive 2; Mismatches
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NAME: Billings, Lucy J.
REGISTRATION NUMBER: 36,749
REFERENCE/DOCKET NUMBER: PF-0179 US
TELECOMMUNICATION INFORMATION:
TELEPHONE: 415-855-0555
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match 76.5%;
Best Local Similarity 75.0%;
Matches 6; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ; ORGANISM: Candida albicans
US-09-248-796A-22159
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APPLICATION NUMBER:
FILING DATE:
Best Local Similarity 75.0
Matches 6; Conservative
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210 IYDGMEHV 217
                                                                                  2 LYDGMEHL 9
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TYPE: PRT
ORGANISM: Human MAGE-A4 peptide
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                                                                                                                                                                                                                                                                                                                                                                                                           FACENCE NO. SELZASI
; GENERAL INFORMATION:
; APPLICANT: VENTER, J. Craig et al.
; APPLICANT: VENTER, J. Craig et al.
; TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF;
; TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF;
; CURRENT APPLICATION NUMBER: US/09/949,016
; CURRENT PILING DATE: 2000-04-14
; PRIOR FILING DATE: 2000-10-20
; PRIOR FILING DATE: 2000-10-03
; PRIOR APPLICATION NUMBER: 60/231,498
; PRIOR FILING DATE: 2000-10-03
; PRIOR FILING DATE: 2000-09-08
; NUMBER OF SEQ ID NOS: 207012
; SEQ ID NOS: 207012
; SEQ ID NO 1171
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                                                                                                                                                                                                        76.5%; Score 39; DB 2; Length 347; 77.8%; Pred. No. 16; 2; Indels ive 0; Mismatches 2; Indels
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18;
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                                                                                                                                                                                                                                                                                                                                                                                Sequence 11171, Application US/09949016
Patent No. 6812339
          TELEX:
INFORMATION FOR SEQ ID NO: 3:
SEQUENCE CHARACTERISTICS:
LENGTH: 347 amino acide
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
IMMEDIATE SOURCE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity 77.0
Best Local Similarity 77.0
                                                                                                                                                                                                        Query Match 76.5
Best Local Similarity 77.8
Matches 7; Conservative
TELEFAX: 415-845-4166
                                                                                                                                                                                                                                                                                                     228 GAYDGEEHL 236
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                                                                                                                                         i LIBRARY: GenBank
; CLONE: 608993
US-08-773-870-3
                                                                                                                                                                                                                                                                                                                                                                      US-09-949-016-11171
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ORGANISM:
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Sequence 28162, Application US/09252991A

TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO PSEUDOMONAS

TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO PSEUDOMONAS

TITLE OF INVENTION: AERUGINOSA FOR DIAGNOSTICS AND THERAPEUTICS

FILE REFERENCE: 107196_136

CURRENT PAPLICATION NUMBER: US/09/252,991A

CURRENT FILING DATE: 1998-02-18

PRIOR FILING DATE: 1998-02-18

PRIOR FILING DATE: 1998-07-27

NUMBER OF SEQ ID NOS: 33142

LENGTH: 494
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APPLICANT: Luiten, Marie-Therese
APPLICANT: Duffour, Marie-Therese
APPLICANT: Demotte, Nathalie
APPLICANT: Demotte, Nathalie
APPLICANT: Demotte, Nathalie
APPLICANT: Cornelis, Guy
APPLICANT: Lurquin, Christophe
APPLICANT: Lurquin, Christophe
APPLICANT: Lurquin, Christophe
APPLICANT: Lurquin, Thierry
APPLICANT: Moon-Falleur, Thierry
APPLICANT: Moon-Falleur, Thierry
APPLICANT: Boon-Falleur, Thierry
APP
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                                                                                                                                                                                                                                                                                                                                       Score 39; DB 4
Pred. No. 18;
0; Mismatches
NUMBER OF SEQ ID NOS: 207012
SOFTWARE: PastSEQ for Windows Version 4.0
SEQ ID NO 11172
LENGTH: 397
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; Sequence 44, Application US/09165863
; Datent No. 6407063
; GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ORGANISM: Pseudomonas aeruginosa
                                                                                                                                                                                                                                                                                                                                            76.5%;
                                                                                                                                                                                                                                                                                                                                                Query Match 76.5
Best Local Similarity 77.8
Matches 7; Conservative
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Best Local Similarity
7; Conserva
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                                                                                                                                                                        ; TYPE: PRT
; ORGANISM: Human
US-09-949-016-11172
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PRIOR FILING DATE: 1998-10-02
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US-09-949-016-9310
                                                                                                                                                                                                                                                                                                                                                                                                                                                             RESULT 14
US-09-949-016-9310
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                                                                                                                                                                                                                                                                                                                         APPLICANT: Chaux, Pascal
APPLICANT: Chaux, Pascal
APPLICANT: Luiten, Rosalie
APPLICANT: Demotter, Nathalie
APPLICANT: Demotter, Nathalie
APPLICANT: Deffour, Maria-Therese
APPLICANT: Traversari, Catla
APPLICANT: Stroobant, Vincent
APPLICANT: Boon-Falleur, Thierry
APPLICANT: Cornelis, Guy R.
APPLICANT: Deon-Falleur, Thierry
APPLICANT: Deon-Falleur, Thierry
APPLICANT: Wan Der Bruggen, Pierre
TITLE OF INVENTION: THOMA ANTIGENS AND CTL CLONES ISOLATED BY A NOVEL
TITLE OF INVENTION: PROCEDURE
TITLE OF INVENTION: PROCEDURE
TITLE OF INVENTION NUMBER: US/09/289,350
CURRENT APPLICATION NUMBER: 1999-04-09
PRIOR APPLICATION NUMBER: 1998-10-02
NUMBER OF SEQ ID NOS: 49
SOFTWARE: Patentin Ver. 2.1
LENGTH: 10
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         APPLICANT: Chaux, Pascal
APPLICANT: Luiten, Rosalie
APPLICANT: Demotre, Nathalie
APPLICANT: Demotre, Nathalie
APPLICANT: Deffour, Marie-Therese
APPLICANT: Traversari, Catia
APPLICANT: Traversari, Catia
APPLICANT: Stroobant, Vincent
APPLICANT: Sconalis, Guy R.
APPLICANT: Boon-Falleur, Thierry
APPLICANT: Wan Der Bruggen, Pierre
APPLICANT: Schultz, Erwin
APPLICANT: Warnler, Guy
TITLE OF INVENTION: TUNCEDURE
TITLE OF INVENTION: TWOCEDURE
FILE REFERENCE: 11727zy
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                                       DB 4; Length 10;
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                                                                              1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         1; Indels
                                74.5%; Score 38; DB 4; 75.0%; Pred. No. 0.49; iive 1; Mismatches
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Pred. No. 0.49;
1; Mismatches
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CURRENT FILING DATE: 2001-04-03
PRIOR APPLICATION NUMBER: 09/165,863
                                                                                                                                                                                                                             RESULT 12
US-09-289-350-44
Sequence 44, Application US/09289350
; Patent No. 651451
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Patent No. 6710172
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              TYPE: PRT
ORGANISM: Human MAGE-A4 peptide
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Best Local Similarity 75.0
Matches 6; Conservative
                                    Query Match 74.5
Best Local Similarity 75.0
Matches 6; Conservative
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1 GVYDGREH 8
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US-09-806-769-44
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US-09-165-863-44
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US-OU-949-016-23-10.

US-OU-949-016-23-10.

US-OU-949-016-23-10.

US-OU-949-016-23-10.

US-OU-949-016-23-10.

US-OU-949-016-23-10.

US-OU-94-016-20.

TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF.

TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF.

TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF.

PILLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF.

UNRENT APPLICATION NUMBER: US/09/949,016

PRIOR FILING DATE: 2000-10-20

PRIOR PILING DATE: 2000-10-03

PRIOR PILING DATE: 2000-10-03

PRIOR PILING DATE: 2000-10-03

PRIOR PILING DATE: 2000-10-03

WUMBER OF SEQ ID NOS: 207012

SOFTWARE: PEASEQ for Windows Vergion 4.0

LENGTH: 312
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Pred. No. 21;
1; Mismatches 2; Indels
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Pred. No. 0.49;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         1; Mismatches
PRIOR APPLICATION NUMBER: 09/289,350
PRIOR FILING DATE: 1999-04-09
PRIOR PILING DATE: 1999-04-02
PRIOR FILING DATE: 2001-04-02
NUMBER OF SEQ ID NOS: 57
SOFTWARE: PATENTIN Ver. 2.1
SEQ ID NO 44
LENGTH: 10
                                                                                                                                                                                                                                                                                                                                                                                              ; TYPE: PRT
; ORGANISM: Human MAGE-A4 peptide
US-09-806-769-44
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            74.5%;
75.0%;
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Best Local Similarity 75.0
Matches 6; Conservative
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Best Local Similarity 66.7
Matches 6; Conservative
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1 GVYDGREH 8
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COMPUTER READABLE FORM:

MEDIUM TYPE: Diskette
COMPUTER: Diskette
COMPUTER: Diskette
COMPUTER: Diskette
COMPUTER: Tab Comparible
COMPUTER: FastSEQ for Windows Version 2.0
SOFTWARE: FaitSEQ for Windows Version 2.0
CURRENT APPLICATION DATA:
FILING DATE: Filed Herewith
FILING DATE: Filed Herewith
FILING DATE: Filed Herewith
FILING DATE: APPLICATION NUMBER:
MARE: Ballings, Lucy J.
REGISTRATION NUMBER: 36,749
REGISTRATION FOR SEQ ID NO: 5:
FELEPHONE: 650-855-0555
INPORMATION FOR SEQ ID NO: 5:
CENTRAIN SEQUENCE: LENGTH: 23,7 anino acids
FYPE: anino acid
STRANDEDNESS: Single
FYPE: Anino acid
STRANDEDNESS: Single
FYPE: LIBRARY: Genbank
FYPE: FYPE
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Search completed: November 15, 2005, 15:45:52 Job time : 1.45238 secs

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Query Match
Best Local S
Matches
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13, Appl
29, Appl
24, Appl
8, Appli
39, Appl
13, Appl
13, Appl
398, App
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Sequence 13, My
Sequence 29, My
Sequence 24, My
Sequence 39, My
Sequence 39, My
Sequence 392, My
Sequence 392, My
Sequence 398, My
Sequence 398, My
                                                                                                                       ; Search time 5.35714 Seconds (without alignments)
702.928 Million cell updates/sec
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Published Applications AA:*

1: /cgn2_6/ptodata/1/pubpaa/VSO7_PUBCOMB.pep:*
2: /cgn2_6/ptodata/1/pubpaa/PCT_REW_PUB.pep:*
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GenCore version 5.1.6
(c) 1993 - 2005 Compugen Ltd.
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US-10-164-121A-13
US-10-164-078A-12
US-10-447-161-29
US-10-651-616-24
US-10-76-51-8
US-10-776-521B-392
US-10-776-521B-393
US-10-776-521B-398
US-10-820-067A-896
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                                                                                                                                                                                                                                                                                                                                                          1867879 segs, 418409474 residues
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   SUMMARIES
                                                                                                                         November 15, 2005, 15:35:58
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Maximum Match 100%
Listing first 45 summaries
                                                                                 - protein search, using sw model
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Gapop 10.0 , Gapext 0.5
                                                                                                                                                                                                    US-09-856-812B-42
51
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Maximum DB seq length: 200000000
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Match Length
                   Copyright
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                                                                                                                                                                                                                                   Perfect score:
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                                                                                                                                                                                                                                                       Sequence:
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Sequence 149, App Sequence 4, Appli Sequence 3202, App Sequence 29, Appli Sequence 29, Appli Sequence 29, Appli Sequence 2919, App Sequence 2919, App Sequence 3041, Appli Sequence 302, Appli Sequence 302, Appli Sequence 17, Appli Sequence 18, Appli Sequence 18, Appli Sequence 27, Appli Sequence 28, Appli Sequence 3170, Appli Sequence 3170, Appli Sequence 27, Appli Appli 27, Appli 27, Appli 27, Appli 27, Appli 27, Appli 27, Appl
# US-10-036-542-84

| US-10-186-812-149
| US-10-186-812-149
| US-10-056-149-4721
| US-10-020-386-32058
| US-10-020-386-32058
| US-10-020-136-3202
| US-10-036-432-29
| US-10-036-432-29
| US-10-0425-114-68250
| US-10-425-114-68250
| US-10-425-114-68250
| US-10-425-115-304865
| US-10-425-115-304870
| US-10-656-884-3
| US-10-656-884-3
| US-10-656-884-3
| US-10-647-115-9
| US-10-164-121A-18
| US-10-165-548A-20
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Sequence 39, Application US/09865548A

Sequence 39, Application US/09865548A

Sequence 39, Application No. US20030096298A1

GENERAL INFORMATION:
APPLICANT: Bear. Ilan
APPLICANT: Admon, Arie
TITLE OF INVENTION: METHOD OF IDENTIFYING PEPTIDES CAPABLE OF BINDING TO MHC MOLECULE
TITLE OF INVENTION: METHOD OF IDENTIFYING PEPTIDES CAPABLE OF BINDING TO MHC MOLECULE
TITLE OF INVENTION: PEPTIDES IDENTIFIED THEREBY AND THEIR USES
TITLE OF INVENTION: WUMBER: US/09/865,548A

CURRENT FILING DATE: 2001-05-16

PRIOR FILING DATE: 2001-05-16

PRIOR FILING DATE: 2001-05-16

NUMBER OF SEQ ID NOS: 204

SEQ ID NOS: 204

SEQ ID NO 39

TENCHALL OF SEC ID NOS: 204

TENCHALL OF SEC ID NOS: 204
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ilarity 100.0%; Pred. No. 1.7e+06;
Conservative 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ) OTHER INFORMATION: synthetic peptide US-09-865-548A-39
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 TYPE: PRT ORGANISM: Artificial Sequence
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9; Conserva
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US-10-651-616-24

US-10-651-616-24

Sequence 24, Application US/10651616

Publication No. US20040253235A1

GENERAL INFORMATION:

APPLICANT: Cytcoure, LLC

APPLICANT: Cytcoure, LLC

APPLICANT: WIRNICK, JAMES T.

TITLE OF INVERTION: METHODS FOR UP-REGULATING ANTIGEN EXPRESSION IN TUMORS

FILE REFERENCE: 027823-0305578

CURRENT APPLICATION NUMBER: US/10/651,616

CURRENT PLINE DATE: 2003-08-29

PRIOR FILING DATE: 2003-08-29

NUMBER OF SEQ ID NOS: 28

SOFTWARE: Patentin version 3.1

SEQ ID NO 24

LENGTH: 9
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GENERAL INFORMATION:
GENERAL INFORMATION:
GENERAL INFORMATION:
GENERAL INFORMATION:
THOUSE OF INVENTION:
METHOD FOR READY-TO-USE, ANTIGEN LOADED OR UNLOADED,
TITLE OF INVENTION: GEYCONSERVED MATURE DENDRITIC CELLS
FILE REFERENCE: 1430/12
CURRENT APPLICATION UNDER: US/10/362,715
CURRENT FILING DATE: 2003-02-24
NUMBER OF SEQ ID NOS: 27
SOFTWARE: PATENTIN VERSION 3.2
SEQ ID NO 8
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                                                                                                                                                                                                                                                    Query Match
100.0%; Score 51; DB 15;
Best Local Similarity 100.0%; Pred. No. 1.7e+06;
Matches 9; Conservative 0; Mismatches 0;
                                                                                                                                                                            ; OTHER INFORMATION: Synthetic Peptide US-10-447-161-29
PRIOR FILING DATE: 2002-05-28
NUMBER OF SEQ ID NOS: 148
SOFTWARE: Patentin version 3.1
SEQ ID NO 29
LENGTH: 9
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                                                                                                                                   ORGANISM: Artificial Sequence FRATURE:
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US-10-362-715-8
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                                                       Sequence 13, Application US/10164121A
Publication No. US20030228308A1
GENERAL INFORMATION:
APPLICANT: Zhang, Xi
APPLICANT: Wan der Bruggen, Pierre
APPLICANT: Traversari, Catra
TITLE OF INVENTION: Isolated Peptides Which Bind to HLA-Cw6 Molecules And Uses There
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        APPLICANT: Schultz, Erwin
APPLICANT: Panichelli, Christophe
APPLICANT: Panichelli, Christophe
APPLICANT: Van der Bruggen, Plerre
APPLICANT: Won der Bruggen, Plerre
TITLE OF INVENTION: Isolated Peptides Which Bind to HLA-B18 Molecules And Uses Therec
FILE REFERENCE: LUD-5756
CURRENT APPLICATION NUMBER: US/10/164,078A
CURRENT FILING DATE: 2002-06-05
NUMBER OF SEQ ID NOS: 35
SEQ ID NO 12
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Publication No. US20040023314A1
GENERAL INFORMATION:
APPLICANT: Wang, Rong-fu
TITLE OF INVENTION: Mutant Fibronectin and Tumor Metastasis
FILE REFRENCE: HO-P02484US1
CURRENT APPLICATION UNMBER: US/10/447,161
CURRENT FILING DATE: 2003-05-28
PRIOR APPLICATION NUMBER: 60/383,530
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100.0%; Pred. No. 1.7e+06;
cive 0; Mismatches 0;
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CURRENT FILING DATE: 2002-08-26
NUMBER OF SEQ ID NOS: 36
SEQ ID NO 13
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Sequence 12, Application US/10164078A Publication No. US20030228325A1 GENERAL INFORMATION:
APPLICANT: Bilsborough, Janine
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Best Local Similarity 100.
Matches 9; Conservative
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Best Local Similarity 100.
Matches 9; Conservative
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ORGANISM: H. sapiens
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ORGANISM: H. sapiens
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US-10-164-078A-12
                     RESULT 2
US-10-164-121A-13
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US-10-447-161-29
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APPLICANT: VAN DER BRUGGEN, Pierre
APPLICANT: BOON-FALLEUR, Thierry
APPLICANT: BOON-FALLEUR, Thierry
APPLICANT: BRECKPOT, Karine
APPLICANT: THIELEMANS, KT.is
TITLE OF INVENTION: ISOLATED PRETIDE WHICH BINDS TO HLA-CW*07 AND USES THEREOF
TITLE OF INVENTION: ISOLATED PRETIDE WHICH BINDS TO HLA-CW*07 AND USES THEREOF
TITLE OF INVENTION: 10041307)
CURRENT APPLICATION NUMBER: US 60/504,874
PRIOR PILING DATE: 2003-09-22
NUMBER OF SEQ ID NOS: 36
SOFTWARE PATENTIN OF SEQ ID NOS: 36
SEQ ID NO 13
LENGTH: 9
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APPLICANT: Fletchner, Jessica
APPLICANT: Prince-Cohane, Kenya
APPLICANT: Prince-Cohane, Kenya
APPLICANT: Bridan
APPLICANT: Balastewicz, Paul
APPLICANT: Barber, Bridan
ITILE OF INVENTION: IMMUNOTHERAPIES
ITILE APPLICATION NUMBER: 00/503,417
PRIOR APPLICATION NUMBER: 60/503,417
PRIOR APPLICATION NUMBER: 60/503,417
PRIOR PILING DATE: 2003-09-16
PRIOR PILING DATE: 2003-09-17
PRIOR PILING DATE: 2003-09-17
PRIOR FILING DATE: 2003-09-17
PRIOR FILING DATE: 2003-09-11
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; TYPE: PRT
; ORGANISM: Artificial Sequence
; PEATURE:
; OTHER INFORMATION: Heat shock protein binding domain
US-10-776-521B-392
                                                                                                                                                                                                                                                                                                                                                  0; Mismatches
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; OTHER INFORMATION: Derived from MAGEA10 US-10-941-150A-13
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ORGANISM: Artificial Sequence
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Best Local Similarity 100.
Matches 9; Conservative
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Best Local Similarity 100.
Matches 9; Conservative
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## Sequence 39, Application US/10705459
## Sequence 39, Application No. US20050053918A1
## Sequence No. US20050053918A1
## Sequence No. US20050053918A1
## Sequence 1. In the Common of th
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APPLICANT: Pletchner, Jessica
APPLICANT: Prince-Cohane, Kenya
APPLICANT: Prince-Cohane, Kenya
APPLICANT: Blusarewicz, Paul
APPLICANT: Andjelic, Sofija
APPLICANT: Barber, Brian
TITLE OF INVENTION: IMMOVED HEAT SHOCK PROTEIN-BASED VACCINES AND
TITLE OF INVENTION: IMMOVED HEAT SHOCK PROTEIN-BASED VACCINES AND
TITLE OF INVENTION: IMMOVED HEAT SHOCK PROTEIN-BASED VACCINES AND
TITLE OF INVENTION: IMMOVED HEAT SHOCK PROTEIN-BASED VACCINES AND
TITLE OF INVENTION: IMMOVER: 004-02-12
PRIOR APPLICATION NUMBER: 60/503,417
PRIOR APPLICATION NUMBER: 60/463,746
PRIOR PILING DATE: 2003-04-18
PRIOR PILING DATE: 2003-04-18
PRIOR PILING DATE: 2003-04-11
PRIOR PELING DATE: 2003-04-11
PRIOR PELING DATE: 2003-04-13
PRIOR PELING DATE: 2003
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100.0%; Score 51; DB 17; Length 9;
Best Local Similarity 100.0%; Pred. No. 1.7e+06;
Matches 9; Conservative 0; Mismatches 0; Indels
                                                  Length 9;
                                                                                                                                    Indels
                                         Query Match 100.0%; Score 51; DB 16; Best Local Similarity 100.0%; Pred. No. 1.7e+06; Matches 9; Conservative 0; Mismatches 0;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ) OTHER INFORMATION: synthetic peptide US-10-705-459-39
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     TYPE: PRT
ORGANISM: Artificial Sequence
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RESULT 14
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US-10-036-542-84

Sequence 84, Application US/10036542

Sequence 84, Application US/10036542

Publication No. US20030083481A1

GENERAL INFORMATION:

APPLICANT: Birse et al.

TITLE OF INVENTION: 25 Human Prostate and Prostate Cancer Associated Proteins

TITLE REFERENCE: PA002P1

CURRENT APPLICATION NUMBER: US/10/036,542

CURRENT FILING DATE: 2002-01-07

PRIOR APPLICATION NUMBER: PCT/US00/19666

PRIOR FILING DATE: 2000-07-20

PRIOR APPLICATION NUMBER: 60/144,972
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US-10-820-067A-896

$ Sequence 696, Application US/10820067A

$ Publication No. US2005214312A1

$ Publication No. US2005214312A1

$ Publication No. US2005214312A1

$ APPLICANT: Pletchner, J.

$ APPLICANT: Parinee-Cohane, K.

$ APPLICANT: Sluaarewicz, P.

$ APPLICANT: Mehta, S.

$ APPLICANT: Barber, B.

$ TITLE OF INVENTION: IMPROVED HEAT SHOCK PROTEIN-BASED TITLE OF INVENTION: VACCIMES AND IMMUNOTHERAPIES FILE OF INVENTION: UMPROVED HEAT SHOCK PROTEIN-BASED TITLE OF INVENTION: UMPROVED HEAT SOUO.067A

$ TITLE OF INVENTION: UMPGER: 60/462,469

$ PRIOR APPLICATION NUMBER: 60/462,469

$ PRIOR APPLICATION NUMBER: 60/463,746

$ PRIOR PILING DATE: 2003-04-18

$ PRIOR PILING DATE: 2003-04-18

$ PRIOR FILING DATE: 2003-04-18

$ NUMBER OF SEQ ID NOS: 926

$ SOFTWARE: FastSEQ for Windows Version 4.0

$ SEQ ID NO 896
NUMBER OF SEQ ID NOS: 419
SOFTWARE: FastSEQ for Windows Version 4.0
SEQ ID NO 398
LENGTH: 20
                                                                                                                                                   FEATURE:
COTHER INFORMATION: Hybrid antigen US-10-776-521B-398
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US-10-820-067A-896
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                                                                                                    TYPE: PRT
ORGANISM: Artificial Sequence
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Matches 9; Conserv
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GENERAL INFUGRATION:

APPLICANT: Mack, David H.

APPLICANT: Aziz, Natasha

APPLICANT: Ess Biotechnology, Inc.

TITLE OF INVENTION: Methods of Diagnosis of Bladder Cancer, Compositions

TITLE OF INVENTION: and Methods of Screening for Modulators of Bladder

TITLE OF INVENTION: and Methods of Screening for Modulators of Bladder

TITLE OF INVENTION: Cancer

FILE REPERENCE: 01801-0023304

FRICK APPLICATION NUMBER: US 60/310,099

PRIOR FILING DATE: 2001-08-03

PRIOR FILING DATE: 2001-08-03

PRIOR PELING DATE: 2001-11-08

PRIOR FILING DATE: 2001-11-08

PRIOR FILING DATE: 2001-11-13

PRIOR FILING DATE: 2001-11-13

PRIOR FILING DATE: 2001-11-13

PRIOR FILING DATE: 2001-11-13

SEQ ID NO 149

LENGTH: 369

"WAMBER OF SEQ ID NOS: 207

SEQ ID NO 149

LENGTH: 369

"WANTER OF SEC ID NOS: 207
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PRIOR APPLICATION NUMBER: 60/148, 681
PRIOR PILING DATE: 1999-08-13
PRIOR PILING DATE: 1999-08-13
PRIOR PILING DATE: 1999-08-17
PRIOR PILING DATE: 1999-06-17
PRIOR APPLICATION NUMBER: 60/158, 004
PRIOR PILING DATE: 1999-10-06
PRIOR PILING DATE: 2000-04-05
NUMBER OF SEQ ID NOS: 157
SOFTWARE: PATCHILING VEY: 2.0
SSQ ID NO 84
LENGTH: 369
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           US-10-188-832-149
. Sequence 149, Application US/10188832
. Publication No. US20040076955A1
. GENERAL INFORMATION:
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; Sequence 4, Application US/10658884
; Publication No. US20050019304A1
; GENERAL INFORMATION:
    APPLICANT: Bandman, Olga
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity 100.
Matches 9; Conservative
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                                                                                                                                                                                                                                                                                                                                                 TYPE: PRT
CORGANISM: Homo sapiens
US-10-036-542-84
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Search completed: November 15, 2005, 15:49:49 Job time: 6.35714 sec8
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254 GLYDGMEHL 262
                                                                                                                                        1 GLYDGMEHL 9
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Sequence 4721, Application US/10756149

Sequence 4721, Application US/10756149

Sequence 4721, Application US/10756149

Sequence 4721, Application US/10756149

Sequence 4721, Natasha

Sequence 4721, Natasha

Septimization Volument and Albert

TITLE OF INVENTION: NOVEL METHODS OF DIAGNOSIS OF METASTATIC CANCER, COMPOSITIONS AND TITLE OF INVENTION: METHODS OF SCREENING FOR MODULATORS OF METASTATIC CANCER

TITLE OF INVENTION: METHODS OF SCREENING FOR MODULATORS OF METASTATIC CANCER

TITLE OF INVENTION: METHODS OF SCREENING FOR MODULATORS OF METASTATIC CANCER

CURRENT APPLICATION NUMBER: US/10/756,149

CURRENT FILING DATE: 2004-01-12

NUMBER OF SEQ ID NOS: 5818

SOFTWARE: PATENT VOLUMENT OF SEQ ID NOS: 5818

SEQ ID NO 4721

LENGTH: 369
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Goli, Surya K.

TITLE OF INVENTION: NOVEL HUMAN MAGE-LIKE PROTEIN
NUMBER OF SEQUENCES: 6
CORRESPONDENCE ADDRESS:
ADDRESSEE: Incyte Pharmaceuticals, Inc.
STREET: 3174 Porter Drive
CITY: Palo Alto
STATE: CA
                                                                                                                                                                                                                                                                                                             COMPUTER READABLE FORM:
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FastSEQ for Windows Version 2.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                CURRENT APPLICATION DATA:

APPLICATION NUMBER: US/10/658,884
FILING DATE: 09-Sep-2003
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US/08/773,870
FILING DATE: 27-DEC-1996
ATTORNEY/AGENT INFORMATION:
NAME: BILINGS, LUC J.
REGISTRATION NUMBER: 36,749
REFERENCE/DOCKET NUMBER: 36,749
TELECOMUNICATION INFORMATION:
TELECOMUNICATION INFORMATION:
TELECOMUNICATION INFORMATION:
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TELECOMUNICATION 110FORMATION:
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; SEQUENCE DESCRIPTION: SEQ ID NO: 4:
US-10-658-884-4
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             INFORMATION FOR SEQ ID NO: 4:
SEQUENCE CHARACTERISTICS:
LENGTH: 369 amino acids
TYPE: amino acidd
STRANDEDNESS: single
TOPOLOGY: linear
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ORGANISM: Homo Sapiens
US-10-756-149-4721
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Best Local Similarity
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Sequence 13, Appl
Sequence 12, Appl
Sequence 29, Appl
Sequence 24, Appl
Sequence 9, Appli
Sequence 39, Appli
Sequence 13, Appl
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                                                                                                                                                                                November 15, 2005, 15:50:00; Search time 163 Seconds (without alignments) 23:102 Million cell updates/sec
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GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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US-10-164-121A-13
US-10-164-024-129
US-10-647-161-29
US-10-651-616-24
US-10-76-51-8
US-10-776-521B-392
US-10-776-521B-392
US-10-149-138-391
US-10-149-138-901
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             1867879 seqs, 418409474 residues
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SUMMARIES
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                                                                                             OM protein - protein search, using sw model
                                                                                                                                                                                                                                                                                                                                                                                                                                                   BLOSUM62
Gapop 10.0 , Gapext 0.5
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Match Length DB
                                                                                                                                                                                                                                                                                                                                                                                      1 GLYDGMEHL 9
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                                                                                                                                                                                              Run on:
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Sequence 2092, Ap Sequence 2681, Ap Sequence 3329, Ap	338,	Sequence 901, App Sequence 2092, Ap	2681,	3329	equence 155,	equence 905,	equence 1078,	equence 2689,	equence 3163	3335	ednence 3638,	equence 3658,	equence 3969,	equence 4144,	equence 905	equence 1078,	equence 2689,	equence 3163,	3335,	edneuce 3638,	3658,	ednence 3969,	4144	equence 155,	٩.	equence 41,	896,	e 896,	896,	9.85
10-149-138-2092 10-149-138-2681 10-149-138-3329	US-10-149-138-338	US-10-149-138-901 US-10-149-138-2092	US-10-149-138-2681	US-10-149-138-3329	US-09-865-548A-155	US-10-149-138-905	US-10-149-138-1078	US-10-149-138-2689	US-10-149-138-3163	US-10-149-138-3335	US-10-149-138-3638	US-10-149-138-3658	US-10-149-138-3969	US-10-149-138-4144	US-10-149-138-905	US-10-149-138-1078	US-10-149-138-2689	US-10-149-138-3163	US-10-149-138-3335	US-10-149-138-3638	US-10-149-138-3658	US-10-149-138-3969	US-10-149-138-4144	US-10-705-459-155	US-10-999-364-4	US-10-273-541-41	US-10-699-088-896	US-10-699-113-896	US-10-699-114-896	US-10-806-924-859
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### ALIGNMENTS

RESULT 1

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Sequence 39, Application US/09865548A

Sequence 39, Application US/09865548A

Sequence 39, Application US/09865548A

Sequence 39, Application US/0966598A1

GENERAL INFORMATION:

APPLICANT: Bear, Ilan

APPLICANT: APPLICANT: Amar

APPLICANT: APPLICANTON: BERTING DATE: 2001-05-16

FILE REFERENCE: 01/22080

CURRENT APPLICATION NUMBER: US/09/865,548A

CURRENT FILING DATE: 2001-05-16

PRIOR APPLICATION NUMBER: US 60/290,958

PRIOR APPLICATION NUMBER: US 60/290,958

NUMBER OF SEQ ID NOS: 204

SOFTWARE: Patentin version 3.1

LEMENTAL OF SEQ ID NOS: 204

SEQ ID NO 39
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      OTHER INFORMATION: synthetic peptide US-09-865-548A-39
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TYPE: PRT
ORGANISM: Artificial Sequence
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     1 GLYDGMEHL 9
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Best Local Similarity
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Matches
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Gaps

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US-10-651-616-24

US-10-651-616-24

Sequence 24, Application US/10651616

Publication No. US20040253235A1

GENERAL INFORMATION:
APPLICANT: CytcCure, LLC
APPLICANT: CytcCure, LLC
APPLICANT: WIRDA, PAUL
FILE REFERENCE: 027823-0305578
CURRENT APPLICATION: WHIPDS FOR UP-REGULATING ANTIGEN EXPRESSION IN TUMORS
FILE REFERENCE: 027823-0305578
CURRENT APPLICATION NUMBER: 08/10/651,616
FRICH RILING DATE: 2003-08-29
NUMBER OF SEQ ID NOS: 28
SOFTWARE: Patentin version 3.1
SEQ ID NO 24

LENGTH: 9

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LENGTH: 9

LENGTH: 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ; OTHER INFORMATION: Description of Artificial Sequence: Homo sapien artificial; OTHER INFORMATION: peptides
US-10-651-616-24
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Sequence 8, Application US/10362715;
Sequence 8, Application US/10362715;
Publication No. US20040253574A1
GENERAL INFORMATION:
GENERAL INFORMATION:
TOTALE OF INVENTION:
TITLE OF INVENTION:
TITLE OF INVENTION:
TITLE OF INVENTION:
TITLE OF INVENTION:
CURRENT APPLICATION NUMBER: US/10/362,715
CURRENT APPLICATION NUMBER: US/10/362,715
CURRENT FILING DATE: 2003-02-24
NUMBER OF SEQ ID NOS: 27
SOFTWARE: Patentin version 3.2
LENGTH: 9
                                                                                                                                                                                                                                                              Length 9;
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100.0%; Score 51; DB 16;
Best Local Similarity 100.0%; Pred. No. 1.7e+06;
Matches 9; Conservative 0; Mismatches 0;
                                                                                                                                                                                                                                                           Query Match 100.0%; Score 51; DB 15; Best Local Similarity 100.0%; Pred. No. 1.7e+06; Matches 9; Conservative 0; Mismatches 0;
                                                                                                                                                                   ; FEATURE:
; OTHER INFORMATION: Synthetic Peptide
US-10-447-161-29
PRIOR FILING DATE: 2002-05-28
NUMBER OF SEQ ID NOS: 148
SOFFWARE: PatentIn version 3.1
SEQ ID NO 29
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        TYPE: PRT
ORGANISM: Artificial Sequence
                                                                                                                       TYPE: PRT ORGANISM: Artificial Sequence
                                                                                                                                                                                                                                                                                                            Conservative
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CRGANISM: Homo sapiens
US-10-362-715-8
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                                                Sequence 13. Application US/10164121A
Publication No. US20030228308A1
GENERAL INFORMATION:
GENERAL INFORMATION:
APPLICANT: Shang, Yi
APPLICANT: Boon, Thierry
APPLICANT: Van der Bruggen, Pierre
APPLICANT: Van der Bruggen, Pierre
APPLICANT: Van der Bruggen, Pierre
TITLE OF INVENTION: 1solated Peptides Which Bind to HLA-Cw6 Molecules And Uses Therec
FILE REFERENCE: LUD-5771
CURRENT APPLICATION NUMBER: US/10/164,121A
CURRENT APPLICATION NUMBER: 2002-08-26
NUMBER OF SEQ ID NOS: 36
SEQ ID NO 13
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      APPLICANT: Schultz, Erwin
APPLICANT: Schultz, Erwin
APPLICANT: Panichelli, Christophe
APPLICANT: Van der Bruggen, Pierre
APPLICANT: Wan der Bruggen, Pierre
APPLICANT: Boon, Thierry
TITLE OF INVENTION: Isolated Peptides Which Bind to HLA-B18 Molecules And Uses Therec
CURRENT APPLICATION NUMBER: US/10/164,078A
CURRENT FILING DATE: 2002-06-05
SEQ ID NOS: 35
LENGTH: 9
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Sequence 29, Application US/10447161
Sequence 29, Application US/10447161
Sequence 29, Application US/10447161
GENERAL INFORMATION:
APPLICANT: Wang, Rong-fu
TITLE OF INVENTION: Mutant Fibronectin and Tumor Metastasis FILE REFERENCE: HO-PO2484US1
CURRENT APPLICATION NUMBER: US/10/447,161
CURRENT FILING DATE: 2003-05-28
PRIOR APPLICATION NUMBER: 60/383,530
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100.0%; Pred. No. 1.7e+06;
tive 0; Mismatches 0;
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APPLICANT: Bilsborough, Janine
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
Best Local Similarity 100.
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Best Local Similarity 100.
Matches 9; Conservative
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ORGANISM: H. sapiens
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                                              US-10-164-121A-13
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Gaps

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Sequence 13, Application US/10941150A

Sequence 13, Application US/2085026881A1

Publication No. US20050226881A1

GENERAL INFORMATION:

APPLICANT: WAN DER BRUGGEN, Pierre

APPLICANT: BOON-PALLEUR, Thierry

APPLICANT: BRECKPOT, Karine

THIELMANS, Kris

TITLE OF INVENTION: ISOLATED WHICH BINDS TO HLA-CW*07 AND USES THEREOF

FILE REFERENCE: LUD-5853-US (10411307)

CURRENT FILING DATE: 2004-09-15

PRIOR PILICATION NUMBER: US/10/941,150A

CURRENT FILING DATE: 2003-09-22

NUMBER OF SEQ ID NOS: 36

SOFTWARE: PatentIn version 3.2

SEQ ID NO 13

LENGTH: 9
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Publication No. US20040018971A1

GENUREL INFORMATION:
GENUREL GENURE, Robert
APPLICANT: Celis, Esteban
APPLICANT: Colis, Esteban
APPLICANT:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Gaps
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                                                                                                                                                                     Length 9;
; TYPE: PRT
; CRGANISM: Artificial Sequence
; FEATURE:
; CTHER INFORMATION: Heat shock protein binding domain
US-10-776-521B-392
                                                                                                                                                            Query Match
Best Local Similarity 100.0%; Pred. No. 1.7e+06;
Matches 9; Conservative 0; Mismatches 0;
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OTHER INFORMATION: Derived from MAGEA10
US-10-941-150A-13
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Best Local Similarity 100.
Matches 9; Conservative
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APPLICANT: Barnea Eilon
APPLICANT: Beer, Ilan
APPLICANT: Admon, Arie
APPLICANT: Admon, Arie
APPLICANT: Buchsbaum, Samuel
APPLICANT: Buchsbaum, Samuel
TITLE OF INVENTION: METHOD OF IDENTIFYING PEPTIDES CAPABLE OF BINDING TO MHC MOLECULE
TITLE OF INVENTION: PEPTIDES IDENTIFIED THEREBY AND THEIR USES
TILE REFERENCE: 2684
CURRENT APPLICATION UNMBER: US/10/705,459
CURRENT FILING DATE: 2003-11-12
SOFTWARE: PALENTI DATE: 2003-11-12
SOFTWARE: PALENTI NOS: 372
SOFTWARE: PALENTI NOS: 372
LENGTH: 9
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                                100.0%; Score 51; DB 16; Length 9; 100.0%; Pred. No. 1.7e+06;
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                                                                                                                                                                                                                                                                                                        Sequence 39, Application US/10705459; Sequence 39, Application US/10705459; Publication No. US20050053918A1; GENERAL INFORMATION
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      TYPE: PRT
ORGANISM: Artificial Sequence
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Matches 9; Conservative
                                                     Best Local Similarity 100
Matches 9; Conservative
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US-10-776-521B-392
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                                   Query Match
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5 GMEHL 9
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### PREADLICANT: Pikes, John
### APPLICANT: State, John
### APPLICANT: Celis, Esteban
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Fublication No. US20040018971A1

FUBLICANT: Fikes, John

APPLICANT: Sette, Alessandro

APPLICANT: Southwood.

APPLICANT: Southwood.

APPLICANT: Clesnit, Robert

APPLICANT: Clesnit, Robert

APPLICANT: Cesteban

APPLICANT: Keogh, Esteban

APPLICANT: Keogh, Esteban

APPLICANT: HER2/neu Using Cellular Immune Responses to

TITLE OF INVENTION: HER2/neu Using Peptide and Nucleic Acid Compositions

FILE REFERENCE: 2060.01440011

CURRENT APPLICATION NUMBER: US/10/149,138
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                                                            FEATURE:
; OTHER INFORMATION: Artificially Synthesized Peptide
US-10-149-138-338
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100.0%; Pred. No. 1.7e+06;
iive 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           US-10-149-138-901
; Sequence 901, Application US/10149138
; Publication No. US20040018971A1
TYPE: PRT
ORGANISM: Artificial Sequence
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ORGANISM: Artificial Sequence
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Best Local Similarity 100.
Matches 5; Conservative
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Best Local Similarity 100.
Matches 5; Conservative
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APPLICANT: Fikes, John
APPLICANT: Sette, Alessandro
APPLICANT: Sette, John
APPLICANT: Sette, John
APPLICANT: Sette, John
APPLICANT: Southwood, Scot
APPLICANT: Southwood, Scot
APPLICANT: Clis, Esteban
APPLICANT: Cesqh. Elissa
TITLE OF INVENTION: HERZ/neu Using Peptide and Nucleic Acid Compositions
TITLE OF INVENTION: HERZ/neu Using Peptide and Nucleic Acid Compositions
TITLE OF INVENTION: HERZ/neu Using Peptide and Nucleic Acid Compositions
TITLE OF INVENTION: HERZ/neu Using Peptide
TITLE OF INVENTION NUMBER: US/10/149,138
CURRENT APPLICATION NUMBER: DCT/USO0/33591
PRIOR PELING DATE: 2000-12-11
PRIOR FILING DATE: 2000-12-11
NUMBER OF SEQ ID NOS: 4641
SOFTWARE: Patentin Version 3.1
SEQ ID NO 2681
LENGTH: 8
LENGTH: 8
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ; OTHER INFORMATION: Artificially Synthesized Peptide US-10-149-138-2681
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100.0%; Pred. No. 1.7e+06;
iive 0; Mismatches 0;
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100.0%; Pred. No. 1.7e+06;
tive 0; Mismatches 0;
PRIOR APPLICATION NUMBER: PCT/US00/33591
PRIOR FILING DATE: 2000-12-11
PRIOR PILING DATE: 1999-12-11
NUMBER OF SEQ ID NOS: 4641
SOFTWARE: Patentin version 3.1
ENGIN 10 2092
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Publication No. US20040018971A1
GENERAL INFORMATION:
APPLICANT: Fikes, John
APPLICANT: Sette, Alessandro
APPLICANT: Southwood, Scott
APPLICANT: Couthwood, Scott
APPLICANT: Chesnut, Robert
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ; Sequence 2681, Application US/10149138
; Publication No. US20040018971A1
; GENERAL INFORMATION:
                                                                                                                                                                                               TYPE: PRT
ORGANISM: Artificial Sequence
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ORGANISM: Artificial Sequence
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Best Local Similarity luv...
5; Conservative
                                                                                                                                                                                                                                                                                                                               Query Match
Best Local Similarity 100.
Matches 5; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                5 GMEHL 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                             GMEHL 5
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APPLICANT: Celis, Esteban
APPLICANT: Keogh, Elissa
TITLE OF INVENTION: Inducing Cellular Immune Responses to
TITLE OF INVENTION: HERZ/neu Using Peptide and Nucleic Acid Compositions
TITLE OF INVENTION: HERZ/neu Us/10/149,138
FILE REFERENCE: 2060.0140001
GURRENT APPLICATION NUMBER: US/10/149,138
CURRENT FILING DATE: 2002-06-10
FRIOR APPLICATION NUMBER: PCT/US00/33591
FRIOR FILING DATE: 1999-12-11
FRIOR APPLICATION NUMBER: US 99/458,299
FRIOR FILING DATE: 1999-12-11
NUMBER OF SEQ ID NOS: 4641
SEQ ID NO 3329
LENGTH: 8
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           APPLICANT: Sette, John
APPLICANT: Southwood, Scott
APPLICANT: Chesnut, Robert
APPLICANT: Rocgh, Elissa
TITLE OF INVENTION: Inducing Cellular Immune Responses to
TITLE OF INVENTION: UMBER: US/10/149,138
FILE REFERENCE: 2002-06-10
CURRENT PILING DATE: 2002-06-10
FRIOR FILING DATE: 1999-12-11
FRIOR FILING DATE: 1999-12-11
NUMBER OF SEQ ID NOS: 4641
SEQ ID NO 338
LENGTH: 8
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ö
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       54.9%; Score 28; DB 16; Length 8; 100.0%; Pred. No. 1.7e+06; rive 0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    54.9%; Score 28; DB 15; Length 8; 100.0%; Pred. No. 1.7e+06; Live 0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ; OTHER INFORMATION: Artificially Synthesized Peptide US-10-149-138-338
                                                                                                                                                                                                                                                                                                                                                                                                                                              ; OTHER INFORMATION: Artificially Synthesized Peptide
US-10-149-138-3329
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Sequence 338, Application US/10149138 Publication No. US20040121946A9 GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                         TYPE: PRT
ORGANISM: Artificial Sequence
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           TYPE: PRT
ORGANISM: Artificial Sequence
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Best Local Similarity 100.
Matches 5; Conservative
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Best Local Similarity 100.
Matches 5; Conservative
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Search completed: November 15, 2005, 15:59:48 Job time : 163 secs

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1

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GENERAL INFORMATION:

APPLICANT: Groteau, Rodney B

APPLICANT: Burke, Charles C

APPLICANT: Burke, Charles C

APPLICANT: Burke, Charles C

APPLICANT: Wildung, Mark R

TITLE OF INVENTION: OF USE

FILE REFERENCE: WGUN14294

CURRENT PILING DATE: 1999-10-18

CURRENT APPLICATION NUMBER: US/09/420,211B

CURRENT PILING DATE: 1999-10-15

EARLIER PILING DATE: 1999-10-15

EARLIER PILING DATE: 1999-10-16

EARLIER PILING DATE: 1997-10-16

NUMBER OF SEQ ID NOS: 15

SEQ ID NO: 2.0

SEQ ID NO: 2.0

SEQ ID NO: 2.0
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                          FILING DATE: 01.748.03

ATTORNEY/AGENT INFORMATION:
NAME: Pabet, Patrea L.
REGIGTRATION NUMBER: 31,284

REFERENCE/DOCKET NUMBER: OWRF 114CIP
TELECOMMUNICATION INFORMATION:
TELEFAK: (404) 815-6508

INFORMATION FOR SEQ ID NO: 22:
SEQUENCE CHARACTERISTICS:
LEGGRATION FOR SEQ ID NO: 22:
SEQUENCE CHARACTERISTICS:
LEGGRATION FOR SEQ ID NO: 22:
SEQUENCE CHARACTERISTICS:
LEGGRATION FOR SEQ ID NO: 22:
SEQUENCE TERES: SINGLE TYPE: amino acids
TYPE: amino acid
STRANDENESS: single
TYPE: amino acid
STRANDENESS: single
TYPE: peptide
HYPOTHETICAL: NO
ANTI-SENSE: NO
ANTI
US 07/648205
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10S-09-420-211-12
1 Sequence 12, Application US/09420211B
1 Patent No. 6303330
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Best Local Similarity 80.0
Matches 4; Conservative
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; ORGANISM: Mentha piperita
US-09-420-211-12
APPLICATION NUMBER:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       2 LYDGMEH 8
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Search completed: November 15, 2005, 15:57:00 Job time: 42 secs

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Substrate Specificity of Protein Kinages
88
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                WESULT 14
US-08-335-198-22

US-08-315-198-22

Sequence 22, Application US/08335198

Patent No. 5637454

GENERAL INFORMATION:
APPLICANT: Harley, John B.
TITLE OF INVENTION: Diseases
NUMBER OF SEQUENCES: 41

CORRESPONDENCES:
ADDRESSEE: Kilpatrick & Cody
STREET: 1100 Peachtree Street, Ste. 2800
CITY: Atlanta
STATE: Georgia
COMPUTR: 100 Peachtree Street, Ste. 2800

STATE: Georgia
COMPUTR: INP PC compatible
COMPUTER READABLE FORM:
BELICATION TREES: PATENT PC-DOS/MS-DOS
CORPUTER: INP PC compatible
COMPUTER: PATENT PEACHIN Release #1.0, Version #1.25
CURRENT APPLICATION NUMBER: US/08/335,198
FILING DATE:
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 24; DB 5; I Pred. No. 4.1e+05;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            PRIOR APPLICATION DATA:

APPLICATION NUMBER: US 08/178,570
FILING DATE: JANUARY 7, 1994
FILING DATE: JANUARY 7, 1994
ATTORNEY/AGENT INFORMATION:
NAME: DECONTI, GIULIO A., Jr.
REGISTRATION NUMBER: 31,503
REFERENCE/DOCKET NUMBER: BBI-004CPPC
TELECOMUNICATION INFORMATION:
TELECAMONICATION INFORMATION:
TELECAX: (617) 227-7400
TELEFAX: (617) 227-5941
INFORMATION FOR SEQ ID NO: 72:
SEQUENCE CHARACTERISTICS:
LENGTH: 9 amino acids
TYPE: amino acid
TYPE: amino acid
                        APPLICANT:
TITLE OF INVENTION: Substrate Specifician WHREN OF SEQUENCES: 86
CORRESPONDENCE ADDRESS:
ADDRESSES: LAHIVE & COCKFIELD STREET, SOUTH SOUTH STREET, SOUTH STREET, SOUTH STREET, SOUTH STREET, SOUTH STREET, SOUTH STREET, SOUTH STREET SOUTH SOUTH STREET SOUTH SOUTH
                                                                                                                                                                                                                                                                                                                                                                                                                                        SOFTWARE: ASCII text
CURRENT APPLICATION DATA:
APPLICATION NUMBER: PCT/US95/00147
FILING DATE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   47.18;
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Best Local Similarity 42.9
Matches 3; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           MOLECULE TYPE: peptide
FRAGMENT TYPE: internal
PCT-US95-00147-72
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      1 GLYDGME 7
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3 GIYEALE 9
GENERAL INFORMATION:
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44.4%; Pred. No. 4.1e+05;
tive 2; Mismatches 3; Indels
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                                                                                                                                 COMPUTER READABLE FORM:

COMPUTER READABLE FORM:

MEDIUM TYPE: Floppy disk

COMPUTER: IBM PC compatible

OPERATING SYSTEM: PC-DOS/MS-DOS/ASCII

SOFTWARE: PatentIn Release #1.0, Version #1.30

CURRENT APPLICATION DATA:

ATLING DATE: Concurrently Herewith

CLASSIFICATION: 514

ATTORNEY/AGENT INFORMATION:

NAME: Kitchell, Barbara S.

REGISTRATION NUMBER: 33,928

REFERENCE/DOCKET NUMBER: UFSC:390/KIT

TELECOMMUNICATION INFORMATION:

TELECOMMUNICATION INFORMATION:

TELEFROM (713) 789-2679
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 24; DB 4; 1
Pred. No. 4.1e+05;
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Sequence 14, Application US/09462453
Fatent No. 6723655
GENERAL INFORMATION:
APPLICANT: BURROWS, SCOTT RENTON
APPLICANT: SHERRITT, MARTINA ALISON
TITLE OF INVENTION: CTL EPITOPES FROM EBV
FILE REFERENCE: FERC.010
CURRENT APPLICATION UNDBER: US/09/462,453
CURRENT FILING DATE: 2000-05-30
NUMBER OF SEQ ID NOS: 33
SOFTWARE: PATENTIN VET: 2.1
SEQ ID NO 14
LENGTH: 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    1; Mismatches
      ADDRESSEE: Arnold, White & Durkee STREET: P.O. Box 4433
                                                                                                                        United States of America
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           RESULT 13
PCT-US95-00147-72
; Sequence 72, Application PC/TUS9500147
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TELEFAX: (713) 789-20.
TELEX: 79-0924
INFORMATION FOR SEQ ID NO: 19:
SEQUENCE CHARACTERISTICS:
LENGTH: 9 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             47.1%;
57.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ; TYPE: PRT
; ORGANISM: Epstein Barr Virus
US-09-462-453-14
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity 44.4
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   TYPE: amino acid
STRANDEDNESS: single
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      MOLECULE TYPE: peptide
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              1 GLYDGMEHL 9
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      1 GLYQGCQ 7
                           STREET: P.O. P.TTY: Houston
                                                                                                   Техав
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                                                                                                                        COUNTRY:
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Gapa ö

1; Indels Length 9;

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47.1%;
                                                                                                                                                                                                                         TYPE: PRT
ORGANISM: Artificial Sequence
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity 71.4%;
Matches 5; Conservative
                                                                                                                                                                                                                                                                                                                                                    Query Match 47.1
Best Local Similarity 42.9
Matches 3; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ; TYPE: PRT
; ORGANISM: Arthrobacter sp.
US-09-503-172A-3
                                                                                                                                                                                                                                                                                                                                                                                                                              1 GLYDGME 7
                                                                                                                                                                                                                                                                                                                                                                                                                                                     |:|: :|
3 GIYEALE 9
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         JS-09-503-172A-3
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US-08-403-459-19
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                                                                                                                                                                                                                                                                                                                                                        Sequence 72, Application US/08178570

Patent No. 552167

GENERAL INFORMATION:
APPLICANT: Lewis C. Cantley
APPLICANT: Shou Song yang
TITLE OF INVENTION: Substrate Specificity of Protein Kinases
NUMBER OF SEQUENCES: 77

CORRESPONDENCE ADDRESS:
                                                                                                                                                                   Length 8;
                                                                                                                                                                                                      1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               47.1%; Score 24; DB 1; Length 9; 42.9%; Pred. No. 4.1e+05; ive 3; Mismatches 1; Indels
                                                                                                                                                             47.1%; Score 24; DB 42.9%; Pred. No. 4.1e tive 3; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           COUNTRY: USA
ZIP: O2109-1875
COMPUTER READBLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: ASCII text
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/178,570
FILING DATE: US/08/178,570
FILECHANICATION INFORMATION:
TELECHANICATION INFORMATION:
TELEFAX: (617) 227-5941
INFORMATION FOR SEQ ID NO: 72:
SEQUENCE CHARACTERISTICS:
LENGTH: 9 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADDRESSEE: LAHIVE & COCKFIELD STREET: 60 STATE STREET, suite 510 CITY: BOSTON STATE: MASSACHUSETTS COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Sequence 72, Application US/08369643A; Patent No. 6004757
                                                                                                                                                               Query Match
Best Local Similarity 42.9
Matches 3; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        MOLECULE TYPE: peptide
; FRAGMENT TYPE: internal
US-08-178-570-72
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INFORMATION FOR SEQ ID NO: SEQUENCE CHARACTERISTICS
                                                                 TOPOLOGY: Incar

MOLECULE TYPE: peptide

FRAGMENT TYPE: internal

PCT-US95-00147-45
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity
                                                     amino acid
                                                                                                                                                                                                                                       1 GLYDGME 7
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2 GIYEALE 8
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3 GIYEALE 9
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US-08-178-570-72
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US-08-369-643-72
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; GENERAL INFORMATION:

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TYPE: PRT
; ORGANISM: Artificial Sequence
; PERATURE:
; OTHER INFORMATION: Description of Artificial Sequence:Enolase (Y43)
US-08-369-643-45
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US-08-369-643-45

Sequence 45, Application US/08369643A

Patent No. 6004757
GENERAL INFORMATION:
APPLICANT: Cantley, Lewis C.
APPLICANT: Songwang, Zhou
TITLE OF INVENTION: Substrate Specificity of Protein Kinases
FILE REFERENCE: CNS-001CP
CURRENT APPLICATION NUMBER: US/08/369,643A
CURRENT APPLICATION NUMBER: US 08/178,570
EARLIER APPLICATION NUMBER: US 08/178,570

SALIER FILING DATE: 1994-01-07
NUMBER OF SEQ ID NOS: 92
SOFTWARE: PatentIn Ver. 2.0
SEQ ID NO 45
LENGTH: 8
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Sequence 45, Application PC/TUS9500147
GENERAL INFORMATION:
GENERAL INFORMATION:
TITLE OF INVENTION: Substrate Specificity of Protein Kinases
NUMBER OF SEQUENCES: 8
CORRESPONDENCE ADDRESS:
ADDRESSER: LAHIVE & COCKFIELD
STREET: 60 STATE STREET, suite 510
        h Similarity 100.0%; Score 24; DB 2; Length 8; Similarity 100.0%; Pred. No. 4.1e+05; 4; Conservative 0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match

47.1%; Score 24; DB 3; Length 8;
Best Local Similarity 42.9%; Pred. No. 4.1e+05;
Matches 3; Conservative 3; Mismatches 1; Indels
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PRIOR APPLICATION DATA:
PRIOR APPLICATION NUMBER: US 08/178,570
PILING DATE: JANUARY 7, 1994
ATTORNEY/AGENT INFORMATION:
NAME: DECOURL, Giulio A., Jr.
REGISTRATION NUMBER: 31,503
REFERENCE/DOCKET NUMBER: BBI-004CPPC
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617) 227-7400
TELEPAX: (617) 227-5941
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           CITY: BOSTON
STATE: MASSACHUSETTS
COUNTY: USA
ZIP: 02109-1875
COMPUTER READABLE FORM:
MEDIUM TYPE: Ploppy disk
COMPUTER: IBM PC compatible
COMPUTER: IBM PC compatible
COMPUTER: IBM PC COMPATIBLE
CURRENT APPLICATION DATA:
APPLICATION NUMBER: PCT/US95/00147
Query Match
Best Local Similarity
Matches 4: Concomm
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2 GIYEALE 8
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                                                                                                                           3 YDGM 6
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Patent No. 5817755
GENERAL INPORMATION:
APPLICANT: Eyre, David R.
APPLICANT: Clemens, J. Daniel
APPLICANT: Clemens, J. Daniel
APPLICANT: Synthetic Peptide Analogs of NTx
ITLE OF INVANTIONS Synthetic Peptide Analogs of NTx
NUMBER OF SEQUENCES: 74
CORRESPONDENCE ADDRESS:
ADDRESSEE: Christensen O'Connor Johnson & Kindness
ADDRESSEE: PLLC
STREET: 1420 Fifth Avenue, Suite 2800
                                                                                                                                                                                                                                                                                                                                                                   47.1%; Score 24; DB 1; Length 8; 100.0%; Pred. No. 4.18+05; tive 0; Mismatches 0; Indels
                                                                                                                                                                                                                                             OTHER INFORMATION: Xaa is glutamine or pyrrolidone of OTHER INFORMATION: carboxylic acid.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ZIP: 98101

ZIP: 98101

COMPUTER ENDABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/807,030
FILING DATE: US/08/807,030
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See OTHER INFORMATION: Xaa is pyroglutamic acid
US-08-08-07-030-11
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FRAGMENT TYPE: N-terminal
                                                                                                                           TOPOLOGY: linear
MOLECULE TYPE: peptide
FRAGMENT TYPE: N-terminal
                                                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity 100.0
....hes 4; Conservative
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                   SEQUENCE CHARACTERISTICS:
LENGTH: 8 amino acide
TYPE: amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               CITY: Seattle
STATE: Washington
COUNTRY: U.S.A.
ZIP: 98101
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                                                                                               STRANDEDNESS
                                                                                                                                                                                                                                     LOCATION: 1
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Gaps

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US-US-446-206B-25

Sequence 25, Application US/08446206B

Patent No. 5750647

GENERAL INFORMATION:

APPLICANT: Eyre, David R

APPLICANT: Clemens, J Daniel

APPLICANT: Ochs, Vincent W

TITLE OF INVENTION: Synthetic Peptide Analogs of NTx

NUMBER OF SEQUENCES: 31

CORRESPONDENCE ADDRESS:

ADDRESSEE: Christensen, O'Connor, Johnson & Kindness PLLC
STREET: 1420 Fifth Avenue
CITY: Seattle

COUNTRY: Washington

COUNTRY: Washington

COUNTRY: U.S.A.

ZIP: WA 9810.

ZIP: WAY 1995

ZIP: WAY 1995
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 24; DB 1; I
Pred. No. 4.1e+05;
3; Mismatches 1;
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PILING DATE:
ATTORNEY DATE:
NAME: Shelton, Dennis K
REGISTRATION NUMBER: 26,997
REGISTRATION NUMBER: WROS18360
TELECOMMUNICATION INFORMATION:
THE PER STATEMENT OF 682 8100
COMPUTER READABLE PORM:
MEDIUM TYEE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: ASCII text
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/178,570
FILING DATE: UNFORMATION:
NAME: DECORT!, Glullo A., Jr.
REGISTRATION NUMBER: 31,503
REFERENCE/DOCKET NUMBER: BBI-004
TELECOMONICATION INPORMATION:
TELECHONE: (617) 227-7400
TELEPHONE: (617) 227-7400
TELEPHONE: (617) 227-5941
JINFORMATION FOR SEQ ID NO: 45:
SEQUENCE CHARACTERISTICS:
LENGTH: 8 amino acids
"WIND AMINO ACIDS ACIDS ACIDS AMINO ACIDS ACIDS ACIDS AMINO ACIDS ACIDS AMINO ACIDS AMI
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Best Local Similarity 42.9
Matches 3; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             TELEFAX: 206 224 0779
TELEX:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ) MOLECULE TYPE: peptide
) FRAGMENT TYPE: internal
US-08-178-570-45
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        TYPE: amino acid
TOPOLOGY: linear
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       1 GLYDGME 7
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |:|: :|
2 GIYEALE 8
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Fatent No. 5522167
GENERAL INFORMATION:
GENERAL INFORMATION:
TITLE OF INVENTION: Substrate Specificity of Protein Kinases
TITLE OF INVENTION: Substrate Specificity of Protein Kinases
NUMBER OF SEQUENCES:
CORRESPONDENCE ADDRESS:
ADDRESSEE: LAHIVE & COCKFIELD
STREET: 60 STATE STREET, suite 510
CITY: BOSTON STATE: NASSACHUSETTS
STATE: NASSACHUSETTS
COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ö
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US-09-116-766-22

Sequence 22, Application US/09116766

Patent No. 596898

GENERAL INFORMATION:

GENERAL INFORMATION:

APPLICANT: Burstein, Yigal

APPLICANT: Trainin, Nathan

APPLICANT: Rycus, Avigail

TITLE OF INVENTION: THF-gamma2 Analogs and Pharmaceutical

TITLE OF INVENTION: Compositions Comprising Them
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               49.0%; Score 25; DB 2; Length 8; 62.5%; Pred. No. 4.1e+05; tive 1; Mismatches 2; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         NUMBER OF SEQUENCES: 28
CORRESPONDENCE ADDRESS:
ADDRESSEE KOME & ASSOCIATES
STREET: 30500 No. 5968898thwestern Hwy., Suite 410
CITY: Farmington Hills
STATE: Michigan
COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          SOFTWARE: Patentin Release #1.0, Version #1.25 CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/116,766
FILING DATE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION DATA:
APPLICATION NUMBER: IL 106214
FILING DATE: 01-JUL-1993
ATTORNEY/AGENT INFORMATION:
NAME: Montgomery, Ilene N.
REGISTRATION NUMBER: 38,972
REFERENCE/DOCKET NUMBER: 2163.00050
TELECOMMUNICATION INFORMATION:
TELEPAX: (248) 539-5050
TELEFAX: (248) 539-5056
INFORMATION FOR SEQ ID NO: 22:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ZIP: 48334
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
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Best Local Similarity 62.5
Matches 5; Conservative
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MOLECULE TYPE: peptide
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1 LEDGPKHL 8
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US-08-178-570-45
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Best Local Similarity 62.5
Matches 5; Conservative
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                                                                                                                                                     November 15, 2005, 15:45:59; Search time 41 Seconds (without alignments) 16.386 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Sequence 22,
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1: /cgn2_6/ptodata/1/iaa/5A_COMB.pep:*
2: /cgn2_6/ptodata/1/iaa/5B_COMB.pep:*
3: /cgn2_6/ptodata/1/iaa/6A_COMB.pep:*
4: /cgn2_6/ptodata/1/iaa/6B_COMB.pep:*
5: /cgn2_6/ptodata/1/iaa/PCTUS_COMB.pep:*
6: /cgn2_6/ptodata/1/iaa/PCTUS_COMB.pep:*
                  GenCore version 5.1.6
(c) 1993 - 2005 Compugen Ltd
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US-09-116-766-22
US-08-446-2068-25
US-08-807-030-11
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PCT-US95-00147-45
US-08-178-570-72
US-08-178-570-72
US-08-178-570-72
US-09-503-172A-3
US-09-420-19
US-09-420-19
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US-08-91-924A-3
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1 GLYDGMEHL 9
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GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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US-09-856-812B-42 51 1 GLYDGMEHL 9 Perfect score: Sequence: ritle:

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

2105692 segs, 386760381 residues Searched: 330156 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 9

Listing first 45 summaries Post-processing: Minimum Match 0% Maximum Match 100%

geneseqp20018:* geneseqp20028:* geneseqp2003as:* geneseqp2003bs:* A_Geneseq_16Dec04:* geneseqp1980s:* geneseqp1990s:* geneseqp2000s:* Database :

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

geneseqp2004s:*

# SUMMARIES

	Description	Aay71487 Human MAG	Aab31323 Exemplary	_	Abj19876 MHC bindi	Adg89586 Class I H	Adil9869 Human HLA	_	Adj58374 Peptide p	Meland	Adre9760 Novel hyb	_	Abg79151 Human MAG	Aay46134 Immunogen	Aay46090 Immunogen	Aay46489 Immunogen	Aay46037 Immunogen		_		_	_	Ade78951 Interleuk	Ade78688 Interleuk	Ade78950 Interleuk	Adh48474 HLA-A*02
SUMMAKIES	ΩĿ	AAY71487	AAB31323	AA017088	ABJ19876	ADG89586	ADI19869	ADI19026	ADJ58374 "	ADL26560	ADR69760	AAY71488	ABG79151	AAY46134	AAY46090	AAY46489	AAY46037	AAY46435	AAG88669	AAG88319	ABJ19990	ADP25980	ADE78951	ADE78688	ADE78950	ADH48474
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# ALIGNMENTS

AAY71487 standard; peptide; 9 AA (first entry) 12-OCT-2000 AAY71487; RESULT 1 AAY71487 

Human MAGE-A10 nonapeptide-1.

MAGE-A10; human; Tumour Rejection Antigen; TRA; Human Leucocyte Antigen; HLA; Major Histocompatibility Complex; MHC; cytolytic T-lymphocyte; CTL; immune response stimulator; prophylaxis; therapy; diagnosis; tumour; cancer; TNF; tumour necrosis factor; vaccine; cytostatic.

Homo sapiens.

WO200032769-A2.

08-JUN-2000

99WO-IB002018. 26-NOV-1999; 98GB-00026143. 27-NOV-1998; (LUDW-) LUDWIG INST CANCER RES.

Boon T; De Plaen E, Brasseur F, Van Pel A, Huang L,

WPI; 2000-412317/35.

Novel polypeptides expressed in tumor cells useful for treating cancers have an ability to complex with a major histocompatibility complex molecule and comprises a specific unbroken amino acid sequence.

Claim 8; Page 36; 80pp; English.

The patent discloses MAGE-A10 and MAGE-A8 polypeptide, nonapeptide and decapeptide sequences, that function as tumour rejection antigens (TRAS). These peptides are capable of forming a complex with major histocompatibility complex (WHC) molecule type HLA-A2.1 (Human Leucocyte Antigen), that are recognised by T-lymphocytes and elicit an immune response from cytolytic T-lymphocytes (TL). They function as an immune response stimulator. Tumour rejection antigens are useful in prophylaxis, therapy and diagnosis of tumours and are effective in controlling or preventing tumour growth. The present peptide sequence is the human MAGE-A10 nonapeptide-1, that corresponds to residues 254-262 of the MAGE-A10 protein. This peptide can serve as a tumour rejection antigen (TRA) and

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Preparation of cryopreserved, mature dendritic cells, useful in vaccines, comprises culturing immature cells on medium containing cocktail of maturation factors, then freezing.
                                                                                                                                       Cryopreserved mature dendritic cell; antigen; vaccine; cytostatic;
                                                                                                             Human mage-A10 protein antigen SEQ ID NO:
                                                                                                                                                    virucide; cancer; hepatitis B virus
                                                                                                                                                                                                                                                                                                                                                                                                                                                            Disclosure; Fig 28; 87pp; German.
                            AAO17088 standard; peptide; 9 AA.
                                                                                                                                                                                                                                                               24-AUG-2001; 2001WO-EP009790.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAB31302-59 represent exemplary antigens which are characteristic of tumours. They can be used to enhance the immune response of vaccines comprising peptides derived from human MAGE-A1 HIA (human leukocyte antigen) class II-binding protein. Human leukocyte antigen) class II-binding protein specification of CD4+T lymphocytes. The MAGE-A1 HIA binding protein stimulate the activity and proliferation of CD4+T lymphocytes. The MAGE-A1 HIA binding protein is useful as a disagnestic agent for diagnosing a disorder characterized by expression of MAGE-A1.

The protein is used for treating a disorder characterized by expression of MAGE-A1 such as cancers e.g. melanomal, squamous cell carcinomas, colorectal carcinomas, osteosarcomas, and lymphocytic leukemias. Peptides derived from the MAGE-A1 HIA binding protein are useful in the production
in combination with adjuvants, can produce vaccines useful for treating a variety of tumours that express MAGE-Al0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Novel MAGE-A1 human leukocyte antigen class II peptides which bind to and are presented to the class II molecules, useful for inducing immune response and treating cancers characterized by expression of MAGE-A1.
                                                                                                                                                                                                                                                                                                        Exemplary antigen characteristic of tumours and derived from MAGE-A10.
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                                                                                                Gaps
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                                                                                                                                                                                                                                                                                                                                  MAGE-A1; HLA; human leukocyte antigen; CD4+ T lymphocyte; cancer;
MAGE-A1 HLA class II-binding protein; vaccine.
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                                                                                             0; Indels
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                                                                  100.0%; Score 51; DB 3; I
100.0%; Pred. No. 1.8e+06;
                                                                                               Mismatches
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                                                                                                                                                                                                                       AAB31323 standard; peptide; 9 AA
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (LUDW-) LUDWIG INST CANCER RES
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                                                   Query Match
Best Local Similarity 100.
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1es 9; Conservative
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                                        Sequence 9 AA;
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The present invention relates to a method for the preparation of readyfor-use, cryopreserved, mature dendritic cells comprising growing immature dendritic cells in a culture medium that includes a 'maturation cockeail' of one or more maturation etimuli and freezing the resulting matured cells in a freezing medium that does not contain heterologous serum. When loaded with antigens, the dendritic cells can be used as vaccines, e.g. against tumours and hepatitis B virus. The present sequence is an antigen described in the invention
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100.0%; Pred. No. 1.8e+06;
ive 0; Mismatches 0;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABJ19876 standard; peptide; 9 AA.
                                                                                                                                                                                                                                                                                           ilarity 100.0%;
Conservative 0
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GLYDGMEHL

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Identifying a cell that differentiates into a metastatic cancer cell, useful for preventing metastatic cancer, comprises identifying a mutated fibronectin in the cell.
                                                                                          The invention comprises a method for identifying a cell that will differentiate into a metastatic cancer cell, the method involves identifying a mutated fibronectin in the cell. The method of the invention is useful for preventing metastatic cancer. The present amino acid sequence represents a Class I HLA-restricted testis cancer antigen.
                                                                       Disclosure; SEQ ID NO 29; 137pp; English
WPI; 2004-035134/03.
                                                                                                                                                                       Sequence 9 AA;
                                                                                                                                                                                                                                                                                                                                                                                      22-APR-2004
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ADI19869
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                                                                                                                                                                                                                                                                    The invention relates to a novel method for identifying peptides originating from a particular cell type, which are capable of binding to major histocompartibility complex (MRC) molecules of a particular haplotype. The method comprises analysing peptides bound to the soluble and secreted form of the MHC molecules of the particular haplotype. The method is useful for identifying peptides for treating an autoimmune disease, such as T or B cell and/or allergic disease or condition, rheumatoid arthritis, or multiple sclerosis, neurodegenerative disorders, e.g. Alzheimer's disease, or diseases associated with inflammation. The sequences of the invention may be used in a gene therapy application. This sequence represents a peptide relating to the method for identifying MHC binding peptides of the invention
                                                                                                                                                                                  Identifying peptides that are capable of binding to major histocompatibility complex (MHG) molecules of a particular haplotype by analyzing peptides bound to the soluble and secreted form of the MHG molecules of the particular haplotype.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    metastatic cancer cell differentiation, mutated fibronectin;
metastatic cancer; class I HLA-restricted; testis; cancer antigen.
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                                                                                                                                      Buchsbaum S;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 9;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                100.0%; Score 51; DB 6; I 100.0%; Pred. No. 1.8e+06;
                                                                                                                                     Beer I, Ziv T, Admon A, Dassau L,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Mismatches
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                                                                                                                                                                                                                                                 Example; Fig 5D; 238pp; English
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                                                                         16-MAY-2001; 2001US-0290958P
29-MAY-2001; 2001US-00865548
                                                 16-MAY-2002; 2002WO-IL000383
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nes 9; Conserv
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 WO200294981-A2
                                                                                                                                                                                                                                                                                                                                                                                                                                         Sequence 9 AA;
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                         28-NOV-2002
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     The present invention relates to a peptide which binds HLA (human leucocyte antigen)-Bl8 to form T-cell epitope. The invention is useful for treating a pathological conditions such as melanoma, lung cancer and head and neck cancer. The present sequence is human HLA-Bl8 binding MAGE-
                                                   Gape
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   HIA; human leucocyte antigen; melanoma; lung cancer; head cancer;
neck cancer; MAGE-3; human.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Panichelli C, Boon-Falleur T;
Length 9;
                                                   Indels
100.0%; Score 51; DB 8; I
100.0%; Pred. No. 1.8e+06;
                                                   0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                    Human HLA-B18 binding MAGE-3 peptide #11.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Claim 22; SEQ ID NO 12; 15pp; English.
                                                                                                                                                                                                                                                                                             ADI19869 standard; peptide; 9 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            05-JUN-2002; 2002US-00164078.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             05-JUN-2002; 2002US-00164078.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Bilaborough J, Schultz E,
                                                                                                                                                                                                                                                                                                                                                                                                    (first entry)
                                                      9; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         SCHULTZ E.
PANICHELLI C.
BOON-FALLEUR T.
BRUGGEN P V D.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   BILSBOROUGH J.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     WPI; 2004-042211/04.
                                                                                                         1 GLYDGMEHL 9
                                                                                                                                                             1 GLYDGMEHL 9
  Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        US2003228325-A1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Sequence 9 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Homo sapiens.
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ADI19026;

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The present invention relates to an isolated peptide which binds to a major histocompatibility complex (MHC) molecule to form a complex that is recognized by a cytolytic T-cell which recognizes and lyses cells presenting complexes of HLA-A2 molecules comprising a sequence of 9 amino acids fully defined in the specification, with the proviso that the peptide is not the peptide of S17 itself. Specifically claimed is an HLA-binding peptide comprising a sequence of 9 amino acids fully defined in the specification. The composition and methods are useful for diagnosing or treating cancer, particularly melanoma. The present sequence
                                                                                                                                                                                                                                                                                                                                                              New isolated SSX-2 and SSX-2-related peptides that bind to human leukocyte antigen (HLA) molecules, useful for diagnosing or treating cancer, particularly melanoma.
                     major histocompatibility complex; MHC; cytolytic T-cell; HLA-A2; Cytostatic; cancer.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   cytostatic, gene therapy; vaccine; cancer; immune response.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         100.0%; Score 51; DB 8; Length 9; 100.0%; Pred. No. 1.8e+06;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Melanoma cell line MAGE gene encoded HLA-A2 peptide.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                   Example 8; SEQ ID NO 34; 20pp; English.
                                                                                                                                                                                                                                           (LUDW-) LUDWIG INST CANCER RES. (TORR-) TORREY PINES INST MOLECULAR
                                                                                                                                                                                                                                                                                              Ayyoub M, Pinilla C;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADL26560 standard; peptide; 9 AA.
                                                                                                                                                                           23-JUL-2003; 2003WO-US023306.
                                                                                                                                                                                                           31-JUL-2002; 2002US-0400076P.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         29-AUG-2002; 2002US-0407492P.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         29-AUG-2003; 2003WO-US027125.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Conservative
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Best Local Similarity
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                                                                                                        WO2004011483-A2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      WO2004019886-A2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Sequence 9 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Homo sapiens
                                                                                                                                        05-FEB-2004
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                                                                                                                                                                                                                                                                                             Valmori D,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      11-MAR-2004
                                                                        Synthetic.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADL26560;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Matches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Treating subject with pathological condition having human leukocyte antigen-Cw6 molecules presented on cell surface by administering peptides to subject for generating immunologically active response against cells.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         The present invention relates to peptides which form immunologically active complexes with MHC molecules. The invention is useful for treating a subject with pathological condition such as cancer which is melanoma by presenting HLA-CW6 molecules on the cell surface. The present sequence is human HLA-CW6 binding MAGE-1 peptide.
                                                                                                                                                                                                                                                                                                                            MHC; major histocompatibility; therapy; cancer; melanoma; HLA-Cw6; human;
                                        Gaps
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     Length 9;
                                   0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             100.0%; Score 51; DB 8; Length 9; 100.0%; Pred. No. 1.8e+06; rive 0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Boon-Falleur T, Bruggen PVD;
   Score 51; DB 8; 1
Pred. No. 1.8e+06;
                                 0; Mismatches
                                                                                                                                                                                                                                                                                         Human HLA-Cw6 binding MAGE-1 peptide #11.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Peptide predicted to bind to HLA-A2 #34.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Example 7; SEQ ID NO 13; 15pp; English.
                                                                                                                                                                                      ADI19026 standard, peptide, 9 AA.
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   100.0%;
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Bost Local Similarity 100.00
Bost Local 9; Conservative
                                                                                                                                                                                                                                                           (first entry)
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                                     9; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Zhang Y, Traversari C,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 (TRAV/) TRAVERSARI C.
(BOON/) BOON-FALLEUR T
(BRUG/) BRUGGEN P V D.
                                                                      σ
                                                                                          GLYDGMEHL 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 1 GLYDGMEHL 9
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Query Match
Best Local Similarity
Matches 9; Conserv
                                                                   1 GLYDGMEHL
                                                                                                                                                                                                                                                                                                                                            MAGE-1; cytostatic
                                                                                                                                                                                                                                                                                                                                                                                                                US2003228308-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ZHANG Y.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Sequence 9 AA;
                                                                                                                                                                                                                                                                                                                                                                                Homo sapiens
                                                                                                                                                                                                                                                           22-APR-2004
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RESULT 8 ADJ58374

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RESULT 10

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The patent discloses MAGE-A10 and MAGE-A8 polypeptide, nonapeptide and decapeptide sequences, that function as tumour rejection antigens (TRAs). These peptides are capable of forming a complex with major histocompatibility complex (MHC) molecule type HLA-A2.1 (Human Leucocyte Antigen), that are recognised by T-lymphocytes and elicit an immune response from cytolytic T-lymphocytes (CTL). They function as an immune response stimulator. Tumour rejection antigens are useful in prophylaxis, therapy and diagnosis of tumours and are effective in controlling or preventing tumour growth. The present peptide sequence is the human MAGE-A8 nonapeptide-1, that corresponds to residues 232.240 of the MAGE-A8
antimicrobial or cytostatic activity. In addition, the invention may prove useful for the production of a vaccine or for gene therapy. The composition and methods disclosed are useful for preventing or treating infectious diseases or cancer. The present sequence is that of a peptide which was used in the exemplification of the invention.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        MAGE-A8; human; Tumour Rejection Antigen; TRA; Human Leucocyte Antigen; Histocompatibility Complex; MHC; cytolytic T-lymphocyte; CTL; immune response stimulator; prophylaxis; therapy; diagnosis; tumour; cancer; TNF; tumour necrosis factor; vaccine; cytostatic.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                protein. This peptide can serve as a tumour rejection antigen (TRA) and in combination with adjuvants, can produce vaccines useful for treating variety of tumours that express MAGE-A8
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Novel polypeptides expressed in tumor cells useful for treating car have an ability to complex with a major histocompatibility complex molecule and comprises a specific unbroken amino acid sequence.
                                                                                                                                                                                       Length 9,
                                                                                                                                                                                                                                  Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  i, Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Boon
                                                                                                                                                                                h Similarity 100.0%; Score 51; DB 8; L Similarity 100.0%; Pred. No. 1.8e+06; 9; Conservative 0; Mismatches 0;
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                                                                                                                                                                                                                                                                                                                                                                                                                                            AAY71488 standard; peptide; 9 AA.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Brasseur
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          (LUDW-) LUDWIG INST CANCER RES
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               98GB-00026143
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human MAGE-A8 nonapeptide-1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (first entry)
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                                                                                                                                                                                                                                                                                                                            GLYDGMEHL
                                                                                                                                                                                                        Local Similarity
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                                                                                                                                            Sequence 9 AA;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               26-NOV-1999;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               27-NOV-1998;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    12-OCT-2000
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                                                                                                                                                                                     Query Match
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Matches
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                                                                                                                                                                                                                                                       response against a tumour cell, comprising administering to a subject with a tumour an amount of IFN-beta receptor agonist and tumour sasociated antigen (TAA). The method is useful for increasing an immune response against a tumour cell. The present sequence is a peptide used in the exemplification of the invention.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                This invention relates to a novel hybrid antigen which comprises at least one antigenic domain of an infectious agent or tumour antigen and a binding domain that non-covalently binds to a heat shock protein. The invention may be useful for the production of compounds with an
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             hybrid antigen; antigenic domain; infectious agent; tumour antigen; binding domain; heat shock protein; antimicrobial; cytostatic; vaccine; gene therapy; infectious disease; cancer.
                                                                                          Increasing an immune response against a tumor cell comprises administering to a subject with a tumor an amount of IFN-beta receptor agonist and tumor associated antigen (TAA).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Slusarewicz P, Andjelic S;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             an antigenic domain and improved heat useful for preventing or treating
                                                                                                                                                                                                                                  The present invention relates to a method of increasing an immune
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                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 9;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                          100.0%; Score 51; DB 8; I 100.0%; Pred. No. 1.8e+06;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Novel hybrid antigen-related peptide #1340.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       0; Mismatches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Example 11; Page 44; 56pp; English.
                                                                                                                                                                                     Example 7; Page 47; 80pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             New hybrid antigens comprising an
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADR69760 standard; peptide; 9 AA.
       Dunn IS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (MOJA-) MOJAVE THERAPEUTICS INC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Fletchner J, Prince-Cohane K,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       domains,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     shock protein-binding domains, infectious diseases or cancer.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     13-FEB-2003; 2003US-0447142P.
11-APR-2003; 2003US-0462469P.
18-APR-2003; 2003US-0463446P.
16-SEP-2003; 2003US-0503417P.
12-PEB-2004; 2004US-00776521.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               13-FEB-2004; 2004WO-US004340
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (first entry)
                                                                                                                                                                                                                                                                                                                                                                                                                                                               Similarity 100.
9; Conservative
       Kurnick JT,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    1 GLYDGMEHL 9
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                                                  WPI; 2004-239114/22.
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                                                                                                                                                                                                                                                                                                                                                                                               Sequence 9 AA;
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treating cancers

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Gaps

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Human MAGE-A10 class I HLA restricted testis cancer antigen peptide #1.
                                                                                                                                                  penetrating peptide; cancer; tumour; melanoma; thymoma; antigen;
                                                                                                                                                          lymphoma; sarcoma; lung cancer; non-Hodgkin's lymphoma; leukaemia; Hodgkin's lymphoma; uterine cancer; cervical cancer; bladder cancer; kidney cancer; adoncarcinoma; breast cancer; prostate cancer; ovarian cancer; pancreatic cancer; epitope; vaccine; dendritic call; tumour infiltrating lymphocyte; TIL; human leukocyte antigen; HLA;
                                                                                                                                                                                                                                                                                                                                                                                                       Novel composition for treating a disease in an animal, comprises an immune effector cell and cell penetrating peptide associated with an antigen or antibody.
                                                                                                                                                                                                                                                                                                                                                                                                                                               Disclosure; Page 14; 61pp; English
                                                                  ABG79151 standard; peptide; 6 AA
                                                                                                                                                                                                                                                                                                                                            (BAYU ) BAYLOR COLLEGE MEDICINE
                                                                                                                                                                                                                                                                                                      15-FEB-2002; 2002WO-US005212.
                                                                                                                                                                                                                                                                                                                         15-FEB-2001; 2001US-0268687P.
                                                                                                          (first entry)
                                                                                                                                                                                                                                                                                                                                                                                   WPI; 2002-627577/67.
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                 GLYDGREH
                                                                                                                                                                                                               cytostatic; human
GLYDGMEH
                                                                                                                                                                                                                                                              WO200264057-A2
                                                                                                                                                                                                                                           Unidentified.
                                                                                                                                                                                                                                  sapiens
                                                                                                          15-NOV-2002
                                                                                                                                                                                                                                                                                  22-AUG-2002
                                                                                       ABG79151;
                                                                                                                                                                                                                                                                                                                                                                Wang R;
                                                                                                                                                                                                                                 Homo
                                                RESULT 12
                                                            ABG7915
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The invention relates to a composition (I) comprising an immune effector cell and a cell penetrating peptide (CPP) associated with an antigen or antibody. Also included are (I) a vaccine comprising (I). CPP associated with an antigen for a disease, by providing (I) and (2) associated with an antigen for disease, by providing (I) and CPP associated with an antigen for disease, by providing (I) and CPP associated with an antigen for disease, by providing the antigens are, for example, tumour antigen derived epitopes recognised by tumour infiltrating lymphocytes (TIL) of HLA (human leukocyte antigen) class I or II. The composition is useful for enhancing immunity in an animal to a disease, by administering a mature dendritic cell comprising CPP associated with an antigen to disease, to the animal, such that following the aministration, animal is protected from disease, where the animal comprises both CD4+ and CD8+ T cells. It is also useful for treating a disease (e.g. cancer, tumour, melanoma, lukmena, lymphoma, sarcoma, lung cancer, non-Hodgin's lymphoma, leukaemia, Hodgin's lymphoma, uterine cancer, cervical cancer, breather animal is further subjected to a cancer and pancearic cancer.

The annimal is further subjected to accer and pancearic cancer.

Sequence 6 AA

radiation, chemotherapy or gene therapy. The administration of (1), preferably dendritic cell is prior to, subsequent to or concurrent with, the cancer treatment. The present sequence is a tumour antigen derived epitope for inclusion in the composition of the invention

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Gaps

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1; Indels

Score 32; DB 2; Length 9; Pred. No. 1.8e+06; 1; Mismatches 1; Indels

Query Match 62.7%; Best Local Similarity 71.4%; Matches 5; Conservative

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AAY45390 to AAY48214 represent specifically claimed immunogenic peptides having a human major histocompatibility complex (MHC) Class I (also known as human leukcoyte antigen (HLA)) binding motif. The immunogenic peptides can bind to a specific HLA allele (i.e. HLA-A subtypes HLA-A2.1, A1, A3.2 or A24.1 or HLA-B or C) and induce a cytotoxic T cell response against the antigen from which the peptide is derived. Cytocoxic T lymphocytes (CLE) which destroy antigen-bearing cells are normally induced by an antigen in the form of a peptide fragment bound to a HLA molecule, rather than the inteat foreign antigen itself, and are particularly important in tumour rejection and in fighting viral infections. The peptides are therefore useful therapeutically to treat or prevent viral infections and cancers in mammals (especially humans) e.g. prostate cancer, hepatitis B and C, AlDS, and renal carcinoma. They can be administered as vaccines to elicit an immune response in individuals susceptible or otherwise at risk conditions. They are also useful diagnostically, and can be used to induce a cytotoxic T cell response, by contacting a cytotoxic T cell with the peptide e.g. to produce TLE ex vivo for infusion back into a produce T cell seponse, by contacting a cytotoxic T cell with the peptide.
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                                                                                                                                                                                                                                                                                                                                Immunogenic peptide having a human leukocyte antigen binding motif #745.
                                                                                                                                                                                                                                                                                                                                                              Human leukocyte antigen; binding; immunogenic; glycoprotein; MHC; HLA; immune response; T cell activation; major histocompatibility complex; cytotoxic T lymphocyte; CTL; tumour rejection; viral infection; cancer; prostate cancer; hepatitis B; hepatitis C; AIDS; renal carcinoma; vaccine; immunisation.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           patient. The polynucleotides encoding the immunogenic peptides are also useful therapeutically and for immunisation as above
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                                        Indela
      Length 6;
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    DB 5; Ler
1.8e+06;
hes 0;
66.7%; Score 34; DB 100.0%; Pred. No. 1.8
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                                                                                                                                                                                                                  AAY46134 standard; peptide; 9 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Claim 1; Page 58; 150pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Sidney J,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          98WO-US005039.
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                                                                                                                                                                                                                                                                                            01-DEC-1999 (first entry)
                                            6; Conservative
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    Query Match
Best Local Similarity
Matches 6; Conserv
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                                                                                1 GLYDGM 6
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Homo sapiens
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              W09945954-A1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              13-MAR-1998;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Synthetic
                                                                                                                                                                                                                                                        AAY46134;
                                                                                                                                                                            RESULT 13
                                                                                                                                                                                              AAY46134
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1 YDGREH

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AAV45390 to AAV48214 represent specifically claimed immunogenic peptides having a human major histocomparibility complex (MHC) Class I (also known as human leukocyte antigen (HLA) binding motif. The immunogenic peptides can bind to a specific HLA allele (i.e. HLA-A subtypes HLA-A2.1, A1, A3.2 or A24.1 or HLA-B or C) and induce a cytocoxic T cell response against the antigen from which the peptide is derived. Cytotoxic T lymphocytes (CTLS) which destroy antigen-bearing cells are normally induced by an antigen in the form of a peptide fragment bound to a HLA molecule, rather than the intact foreign antigen itself, and are particularly important in tumour rejection and in fighting viral infections. The peptides are cancers in mammals (especially to treat or prevent viral infections and cancers in mammals (especially when a pentile sections and cancers in mammals (especially when a definistered as vaccines to elicit an immune response in individuals susceptible or otherwise at risk
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        of viral infection or caneer, or used to treat chronic or acute conditions. They are also useful diagnostically, and can be used to induce a cytotoxic T cell response, by contacting a cytotoxic T cell with the peptide e.g. to produce CTLS ex vivo for infusion back into a patient. The polynucleotides encoding the immunogenic peptides are also useful therapeutically and for immunisation as above
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 New immunogenic peptides with HLA binding motif, useful in treatment and diagnosis of cancers and viral diseases.
                                                                                                                                                                                                                                                    Immunogenic peptide having a human leukocyte antigen binding motif #701.
                                                                                                                                                                                                                                                                                       Human leukocyte antigen; binding; immunogenic; glycoprotein; MHC; HLA; immune response; T cell activation; major histocompatibility complex; cytotoxic T lymphocyte; CTL; tumour rejection; viral infection; cancer; prostate cancer; hepatitis B; hepatitis C; AIDS; renal carcinoma;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Southwood S;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      60.8%; Score 31; DB 2; Length 9; 83.3%; Pred. No. 1.8e+06; ive 0; Mismatches 1; Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Sette A, Kubo RT, Sidney J, Celis E,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Claim 1; Page 56; 150pp; English.
                                                                                                                                    AAY46090 standard; peptide; 9 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    98WO-US005039.
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                                                                                                                                                                                                             (first entry)
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Best Local Similarity 83.3.
                                                                                                                                                                                                                                                                                                                                                                      immunisation
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             WPI; 1999-551214/46.
2 LYDGMEH 8
                      Sequence 9 AA;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                        W09945954-A1.
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                                                                                                                                                                                                                                                                                                                                                                                                               Synthetic
                                                                                                                                                                       AAY46090;
                                                                                                                                                                                                                                                                                                                                                                          vaccine;
                                                                                             RESULT 14
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AAY45390 to AAY48214 represent specifically claimed immunogenic peptides having a human major histocompatibility complex (MHC) Class I (also known as human leukocyte antigen (HLA)) binding motif. The immunogenic peptides can bind to a specific HLA allele (i.e. HLA-A subtypes HLA-A2.1, A1, A3.2 or A24.1 or HLA-B or C) and induce a cytotoxic T cell response against the antigen from which the peptide is derived. Cytotoxic T lymphocytes (CTLS) which destroy antigen-bearing cells are normally induced by an antigen from of a peptide fragment bound to a HLA molecule, rather antigen in the form of a peptide fragment bound to a HLA molecule, rather than the intact foreign antigen from in fraction in factions. The peptides are cumour rejection and in fighting viral infections. The peptides are therefore useful therapeutically to treat or prevent viral infections and cenal carcinoma. They can be administered as vaccines to and c. AlDS, and renal antidividuals succeptible or otherwise at risk of viral infection or cancer, or used to treat chronic or acute conditions. They are also useful diagnostically, and can be used to induce a cytotoxic T cell with the peptide e.g. to produce CTLs ex vivo for infusion back into a patient. The polynucleotides encoding the immunogenic peptides are also patient. The polynucleotides encoding the immunogenic peptides are also
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         New immunogenic peptides with HLA binding motif, useful in treatment and diagnosis of cancers and viral diseases.
                                                                                                                           Immunogenic peptide having a human leukocyte antigen binding motif #1100
                                                                                                                                                    Human leukocyte antigen; binding; immunogenic; glycoprotein; MHC; HLA; immune response; T cell activation; major histocompatibility complex; cytotoxic T lymphocyte; CTL; tumour rejection; viral infection; cancer; prostate cancer; hepatitis B; hepatitis C; AIDS; renal carcinoma; vaccine; immunisation.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Grey HM,
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   134.9%; Score 28; DB ilarity 100.0%; Pred. No. 1.6 Conservative 0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Celis E,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Claim 1; Page 74; 150pp; English.
               AAY46489 standard; peptide; 9 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Sidney J,
                                                                                                                                                                                                                                                                                                                                                                                                                                                        98WO-US005039
                                                                                       01-DEC-1999 (first entry)
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les 5; Conserv
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                                                                                                                                                                                                                                                                                                                                                                                                                    13-MAR-1998;
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                                                                                                                                                                                                                                                                                                     Ното варіеля
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                                                   AAY46489;
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Matches
AAY46489
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Gaps

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3 YDGMEH 8

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GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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OM protein - protein search, using sw model

November 15, 2005, 15:44:54 ; Search time 38 Seconds (without alignments) 22.788 Million cell updates/sec Run on:

Title: Perfect score:

US-09-856-812B-42 51 1 GLYDGMEHL 9 Sequence: BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

283416 segs, 96216763 residues Searched:

791 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 9

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Database :

PIR 79:*
1: pirl:*
2: pir2:*
3: pir3:*
4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

<b></b>	Description	enamelin i - bovin	fibrinopeptide B -	~	chlorophyll a/b-bi	macrophage chemota	aspartate transami	fibrinopeptide B -	glycine cleavage s	thymic humoral fac	whey glycoprotein	endosperm protein,	Ig heavy chain CDR	purine	Ig heavy chain CRD	peptidyl-dipeptida	T-cell receptor be	hypothetical prote	158K exoantigen -	T-cell receptor be	urine glycopeptide	neuropeptide calla	formylglycinamide	protein QA300040 -	phosphatidylethano	-phoe	inulinase (EC 3.2.	sperm-activating p	perm-activ	fibrinogen beta ch
SUMMARIES																														
SCIM	£	510784	E28854	F28854	PW0002	A37027	A11483	D28854	PS0253	A28719	PC4392	870334	PT0225	A56029	PT0240	JN0861	PT0589	T14906	B33099	PT0725	XGHUEU	E47393	A12016	PA0032	PN0043	S11078	PT0030	E60588	G60588	C24180
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Ig heavy chain CRD Ig heavy chain V r late Gl-69 protein dextransucrase (EC gamma subunit of P Ig heavy chain CRD Leu-enkephalin - b Met-enkephalin - b T-cell receptor be biotin B - Citroba T-cell receptor be T-cell recept	
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## ALIGNMENTS

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enamelin i - bovine (fragment)
CiSpecies: Bos primigenius taurus (cattle)
CiSpecies: Bos primigenius taurus (cattle)
CiDate: 19-Mar-1997 #sequence_revision 21-Nov-1998 #text_change 09-Jul-2004
CiAccession: 510784
RiStrawich, E.; Glimcher, M.J.
RiStrawich, E.; Glimcher, M.J.
A;Title: Tooth 'enamelins' identified mainly as serum proteins. Major 'enamelin' is albû A;Reference number: 510780, MUID:90336641; PMID:2379503
A;Accession: 510784
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                                                                                                                                                                                                                                                                                                                   A,Molecule type: protein
A,Residues: 1-9 <STR>
A,Cross-references: UNIPROT:Q7M2M7
C;Keywords: enamel; phosphoprotein
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RESULT 2

E28654

E1brinopeptide B - hamadryas baboon

C;Species: Paplo hamadryas baboon)

C;Species: Paplo hamadryas (hamadryas baboon)

C;Species: Paplo hamadryas (hamadryas baboon)

C;Species: Paplo hamadryas (lamadryas baboon)

C;Baccession: E28854

B;Nakamura, S.; Takanaka, O.; Takahashi, K.

J. Biochem. 94, 1973-1978, 1983

A;Title: Fibrinopeptides A and B of baboons (Paplo anubis, Paplo hamadryas, and Theropit, A;Hocession: E28854

A;Molecule type: protein

A;Molecule type: protein

A;Molecule type: Powdka

A;Residues: 1-9 <NAKs

A;Cross-references: UNIPROT:P19343

C;Superfamily: fibrinogen beta chain; fibrinogen beta/gamma homology; fibrinogen disulfi

Gaps ö Query Match 35.3%; Score 18; DB 2; Length 9; Best Local Similarity 60.0%; Pred. No. 2.8e+05; Matches 3; Conservative 1; Mismatches 1; Indels

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||: | 4 GLFHG 8 1 GLYDG 5

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RESULT 3 F28854

8 셤

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aspareate transaminase (EC 2.6.1.1), mitochondrial - sheep (fragment)
N;Alternate names: aspareate aminotransferase, mitochondrial
S;Species: Ovientalis aries, Ovis ammon aries (domestic sheep)
C;Date: 05-Jun-1987 #sequence_revision 05-Jun-1987 #text_change 20-Aug-1999
C;Accession: Al1483
E;Campos-Cavieres, M.; Milstein, C.P.
Biochem. J. 147, 275-281, 1975
A;Title: The sequences of the coenzyme-binding peptide in the cytoplasmic and the mitoch
A;Reference number: Al1483; MUID:76039441; PMID:1180894
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          fibrinopeptide B - olive baboon
C;Species: Papio anubis, Papio hamadryas anubis (olive baboon)
C;Species: Papio anubis, Papio hamadryas anubis (olive baboon)
C;Accession: D28854
R;Nakamura, S.; Takenaka, O.; Takahashi, K.
R;Nakamura, S.; Takenaka, O.; Takahashi, K.
A;Title: Fibrinopeptides A and B of baboons (Papio anubis, Papio hamadryas, and Theropiti A;Reference number: A91973; WUID:84161822; PMID:6423621
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   A;Wolecule type: protein
A;Residues: 1-9 «NAK»
A;Cross-references: UNIPROT:P19344
C;Superfamily: fibrinogen beta chain; fibrinogen beta/gamma homology; fibrinogen disulfi
                                                                                                                                                                                                                                                                                                                                                                                                                               A;Molecule type: protein
A;Residues: 1-7 <CAM>
A;Experimental source: liver
C;Keywords: aminotransferase; mitochondrion; phosphoprotein; pyridoxal phosphate
F;2/Binding site: pyridoxal phosphate (Lys) (covalent) #status experimental
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Gaps
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Best Local Similarity 100.0
Matches 3; Conservative
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Best Local Similarity 60.v<sup>,</sup>
Local 3; Conservative
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A,Reference number: PS0206
A,Accession: PS0253
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4 GLFRG 8
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fibrinopeptide B - gelada baboon
C; Species: Theropithecus gelada (gelada baboon)
C; Species: Theropithecus gelada (gelada baboon)
C; Species: 19-May-1989 #sequence_revision 19-May-1989 #text_change 09-Jul-2004
C; Accession: F28654
R; Nakamura, S:; Takenaka, O.; Takahashi, K.
J. Blochem. 94, 1973-1978, 1983
A; Title: Fibrinopeptides A and B of baboons (Papio anubis, Papio hamadryas, and Theropit A; Reference number: A91973; MUID:84161822; PMID:6423621
A; Accession: F28854
A; MolColle type: protein
A; Residues: 1-9 < NAK.
A; Residues: 1-9 < NAK.
A; Cross-references: UNIPROT:P19342
C; Superfamily: fibrinogen beta chain; fibrinogen beta/gamma homology; fibrinogen disulfi
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        chlorophyll a/b-binding protein 31K - green alga (Dunaliella tertiolecta) (fragment)
NyAlternate names: photosystem II light-harvesting chlorophyll 31K protein
C;Species: Dunaliella tertiolecte
C;Date: 24-Nov-1999 #sequence_revision 24-Nov-1999 #text_change 24-Nov-1999
C;Accession: PW0002
R;LaRoche, J.; Bennett, J.; Falkowski, P.G.
Gene 95, 165-171, 1990
A;Title: Characterization of a cDNA encoding for the 28.5-kDa LHCII apoprotein from the A;Title: Characterization of a cDNA encoding for the 28.5-kDa LHCII apoprotein from the A;Mclecule type: protein
A;Mclecule type: protein
A;Mclecule type: Drotein
A;Mcseddues: 1-9 <LAR>
C;Superfamily: chlorophyll a/b-binding protein
C;Keywords: chloroplast; grana; light-harvesting complex; membrane adhesion; membrane pr
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A37027
Minacrophage chemotactic factor - human (fragment)
NiAlternate names: T-cell hybridoma D6-18 protein
C,Species: Homo sapiens (man)
C,Species: Homo sapiens (man)
C,Species: Homo sapiens (man)
C,Accession: 33-7027
R;YOShizuka, N; YOShimura, M.; TBuchiya, S.; Okamoto, K.; Kobayashi, Y.; Osawa, T.
Cell. Immunol. 123, 212-225, 1989
A;Yitle: Macrophage chemotactic factor (MCF) produced by a human T cell hybridoma clone.
A;Reference number: A37027; MUID:89376581; PMID:2505934
A;Accession: A37027
A;Accession: Cell. Immunary
A;Residues: preliminary
A;Wolecule type: protein
A;Residues: 1-9 <YOS>
A;Cross-references: UNIPROT:Q7M4S2
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Pred. No. 2.8e+05;
                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match 35.3%; Score 18; DB 2; Length 9; Best Local Similarity 60.0%; Pred. No. 2.8e+05; Matches 3; Conservative 1; Mismatches 1; Indel8
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Pred. No. 2.8e+05;
1; Mismatches 1; Indels
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Best Local Similarity 60.0%;
Matches 3; Conservative 1
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Best Local Similarity 57.14
Matches 4; Conservative
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C'Species: Homo sapiens (man)
C'Dacessers - Coper-1993 #sequence_revision 30-Sep-1993 #text_change 16-Aug-1996
C'Dacession: P70225
R'Yamada, M.; Wasserman, R.; Reichard, B.A.; Shane, S.; Caton, A.J.; Rovera, G.
J. Exp. Med. 173, 395-407, 1991
A;ritle: Preferential utilization of specific immunoglobulin heavy chain diversity and jet.
A;Reference number: P70222; MUID:91108337; PMID:1899102
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C;Species: Mus musculus (house mouse)
C;Species: Mus musculus (house mouse)
C;Accession: A56029
R;Roy, R.; Brooks, C.; Mitra, S.
Biochemistry 33, 15131-15140, 1994
A;Title: Purification and biochemical characterization of recombinant N-methylpurine-DNA
A;Reference number: A56029; MUID:95092772; PMID:7999773
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C; Species: Homo sapiens (man)
C; Species: Homo sapiens (man)
C; Species: 30-Sep-1993 #sequence_revision 30-Sep-1993 #text_change 16-Aug-1996
C; Accession: PTO240
R; Yamada, M: Wasserman, R.; Reichard, B.A.; Shane, S.; Caton, A.J.; Rovera, G.
J. Exp. Med. 173, 395-407, 1991
A; Title: Preferential utilization of specific immunoglobulin heavy chain diversity and jc
A; Reference number: PTO222; MUID:91108337; PMID:1899102
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                                                                                                                                                                                                                                   ig heavy chain CDR3 region (clone 1-103) - human (fragment)
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A,Molecule type: DNA
A,Residues: 1-9 < YAAA
A,Experimental source: B lymphocyte
C,Keywords: heterotetramer; immunoglobulin
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A;Status: preliminary
A;Molecule type: protein
A;Residues: 1-9 <ROY>
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2 MFGGM 6
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                                   7 YEG
3 YDG
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C;Species: Bos primigenius taurus (cattle)
C;Species: 10-Nov-1997 #sequence_revision 10-Nov-1997 #text_change 07-May-1999
C;Accession: PC4392
R;Hwangbo, S.; Azuma, N.; Kurisaki, J.; Kanno, C.
Biosci. Biotechnol. Biochem. 61, 1568-1574, 1997
A;Title: Purification and characterization of novel whey glycoprotein WGP-88 which binds A;Reference number: PC4392; MUID:97480944; PMID:9339560
A;Accession: PC4392
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Cispecies: Secale cereale (rye)
Cipate: 19-Mar-1998 #sequence_revision 17-Apr-1998 #text_change 17-Apr-1998
CiAccession: 870334
RiRocher, A.; Calero, M.; Soriano, F.; Mendez, E.
Biochim. Biophys. Acta 1955, 13-22, 1996
A;Title: Identification of major rye secalins as coeliac immunoreactive proteins.
A;Reference number: 870327; MUID:96283789; PMID:8679669
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A;Residues: 1-6 <HWA-7
A;Experimental source: milk
C;Comment: This protein binds to the monoclonal antibody to PAS-4 glycoprotein.
                                                                                                                                                                                                                                                               Cispecies: Bos primigenius taurus (cattle)
Cipate: 30-Sep-1989 #sequence_revision 30-Sep-1989 #text_change 09-Jul-2004
Cipate: 30-Sep-1989 #sequence_revision 30-Sep-1989 #text_change 09-Jul-2004
Cipatesion: A28719
Riburstein, Y.; Buchner, V.; Pecht, M.; Trainin, N.
Biochemistry 27, 4066-4071, 1988
A;Title: Thymic humoral factor gamma-2: purification and amino acid sequence
A;Reference number: A28719; MUID:88326920; PMID:3261994
A;Reference number: protein
A;Residues: 1-8 < EURx-A;Residues: 1-8 < EURx
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Search completed: November 15, 2005, 15:56:13 Job time : 40 secs

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GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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Bos taurus (Bovine).

Bukaryota, Metazoa, Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovidae;

Bovinae; Bos.

NCB_TaxID=9913;
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Mammalia, Butheria, Primates, Catarrhini, Cercopithecidae,
Cercopithecinae, Papio.
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"Tooth 'enamelins' identified mainly as serum proteins. Major 'enamelin' is albumin.";
Eur. J. Blochem. 191:47-56(1990).
PIR; S10784; S10784.
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01-NOV-1990 (Rel. 16, Last sequence update)
05-JUL-2004 (Rel. 44, Last annotation update)
Fibrinogen beta chain [Contains: Fibrinopeptide B] (Fragment)
                                                                                     Score 20; DB 2; Length 9; Pred. No. 1.6e+06;
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 1 1 1 2 9 9 9 9 1047 MW;
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-1- FTM: Conversion of fibringen to fibrin is triggered by thrombin, which cleaves fibrinoperides A and B from alpha and beta chains and thus exposes the N-terminal polymerization sites responsible for the formation of the soft clot.

FIR: E28854; E28854.

InterPro; IPR002181; Fibrinogen C. PROSITE; PS00514; FIBRIN AG_C_DÖMAIN; PARTIAL.

Blood coagulation; Direct profein sequencing; Plasma.
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Mammalia, Butheria, Primates, Catarrhini, Cercopithecidae,
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01-NOV-1990 (Rel. 16, Last sequence update)
05-UUL-2004 (Rel. 44, Last amotation update)
Fibrinogen beta chain [Contains: Fibrinopeptide B] (Fragment)
                                                                                                                                                                                                                                                                                                                                                                          35.3%; Score 18; DB 1; Length 9; 60.0%; Pred. No. 1.6e+06; tive 1; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    35.3%; Score 18; DB 1; Length 9; 60.0%; Pred. No. 1.6e+06; tive 1; Mismatches 1; Indels
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                                                                                                                                                                                                                                                                                                                           DDFE71E9C7287B06 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     9 AA; 977 MW; DDFE7879C7287B06 CRC64;
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AA
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NCBI_TaxID=9565;
                                                                                                                                                                                                                                                                                                                        9 AA; 1057 MW;
                                                                                                                                                                                                                                                                                                                                                                            Query Match 35.3
Best Local Similarity 60.0
Matches 3; Conservative
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Matches 3; Conserv
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Pubmeda15019625; DOI=10.1016/j.ympev.2003.08.002; Pubmeda15019625; DOI=10.1016/j.ympev.2003.08.002; Paton A., Springate D.A., Sudde S., Otieno D., Grayer R., Harley M.M., Willis P., Simonds M.S.J., Powell M.P., Savolainen V.; Phylogeny and evolution of basils and allies (Ocimeae, Labiatae) based on three plastid DNA regions."; Mol. Phylogenet. Evol. 31:277-229(2004). EMBL; AJ505403; CAD45523.1; EMBL; AJ505403; CAD45523.1; COM5723 (Ocimeae, IEA).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          MEDLINE=84161822; PubMed=6423621;
Nakamura S., Takenaka O., Takahashi K.;
Nakamura S., Takenaka O., Takahashi K.;
Istinopeptides A and B of baboons (Papio anubis, Papio hamadryas, and Trinopeptides A and B of baboons (Papio and Sequences and evolutionary rates and a molecular phylogeny for the baboons.";
J. Blochem. 94:1973-1978(1983).
- I- FUNCTION: Fibrinogen has a double function: yielding monomers that polymerize into fibrin and acting as a cofactor in platelet
                                                                                                              Bukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta; Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; asterids; lamidales; Lamiaceae; Prostantheroideae; Westringleae; Prostantheroideae; Westringleae; Prostanthara.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    -i-SUBUNIT: Hexamer containing 2 sets of 3 nonidentical chains (alpha, beta and gamma), linked to each other by disulfide bonds.
-i-PTM: Conversion of fibrinogen to fibrin is triggered by thrombin, which cleaves fibrinopeptides A and B from alpha and beta chains, and thus exposes the N-terminal polymerization sites responsible for the formation of the soft clot.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi,
Mammalia, Butheria, Primates, Catarrhini, Cercopithecidae,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       01-NOV-1990 (Rel. 16, Created)
01-NOV-1990 (Rel. 16, Last sequence update)
01-NUT-1990 (Rel. 44, Last annotation update)
Fibrinogen beta chain [Contains: Fibrinopeptide B] (Fragment).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 17; DB 2; Length 8; Pred. No. 1.6e+06; 1; Mismatches 1; Indels
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PROSITE; PS00514; FIBRIN AG C DÖMAIN; PARTIAL.
Blood coagulation; Direct protein sequencing; Plasma.
PERTIDE 9 Fibrinopeptide B.
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05-JUL-2004 (TrEMBLrel. 27, Last sequence update) 05-JUL-2004 (TrEMBLrel. 27, Last annotation update) Ribosomal protein (Fragment).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        8 AA; 838 MW; C821F2C058786415 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    9 AA.
                                                                             Prostanthera nivea (snowy mintbush).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 33.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Papio anubis (Olive baboon)
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NCBI_TaxID=9555;
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                 Ribosomal protein.
                                                                                                                                                                                                                                   SEQUENCE FROM N.A.
                                                                                                                                                                                                NCBI_TaxID=38863;
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                                                                                                  Chloroplast
                                                             Name=rps16;
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P19344;
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NON TER
SEQUENCE
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Matches
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Raftery M.J., Bradford A.M., Bowie J.H., Wallace J.C., Tyler M.J.;
"Peptides from Australian frogs. The structure of the dynastins from the banjo frogs Limnodynastes interioras, Limnodynastes dumerilli and Limnodynastes terracreginse.";
Aust. J. Chem. 46:833-842(1993).
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Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Amphibia; Batrachia; Anura; Neobatrachia; Hyloidea; Myobatrachidae;
                                                                                                                                                                                                                                 MEDLINE=89376581; PubMed=2505934;
Yoshizuka N., Yoshimura M., Tsuchiya S., Okamoto K., Kobayashi Y.,
                                                                                                                      Bukaryota, Metazoa; Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo.
NCBI_TaxID=9606;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          -i- MASS SPECTROMETRY: WW=729; WETHOD=FAB; RANGE=1-8; NOTE=Ref.1.
Direct protein sequencing.
SEQUENCE 8 AA; 729 MW; 7C28772865B72728 CRC64;
                                                                                                                                                                                                                                                                                          "Macrophage chemotactic factor (MCF) produced by a human T cell hybridoma clone."; Cell. Immunol. 123:212-225(1989).
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Pred. No. 1.6e+06;
0; Mismatches 3; Indels
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Pred. No. 1.6e+06;
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                                                         Last sequence update)
Last annotation update)
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29-MAR-2004 (Rel. 43, Last sequence update)
05-JUL-2004 (Rel. 44, Last annotation update)
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  9 AA.
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05-JUL-2004 (TrEMBLrel. 27, Created)
                                        Created)
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NCBI_TaxID=30362;
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50.0%;
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                                    01-MAR-2004 (TrEMBLrel. 26, 01-MAR-2004 (TrEMBLrel. 26, 01-MAR-2004 (TrEMBLrel. 26,
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9 AA; 1048 MW;
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Best Local Similarity 50.0
Local 3; Conservative
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  PRELIMINARY:
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                                                                                                                    Homo sapiens (Human)
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Best Local Similarity
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P82079:
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Q712A6;
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                                                                                                                                                                                                                                                                                                                                                                                                Eukaryota, Viridiplantae, Streptophyta, Embryophyta, Tracheophyta; Spekaryota, Magnollophyta; eudicotyledone; core eudicots; asterids; lamiids; Lamiales; Lamiaceae; Nepetoideae; Ocimeae; Alvesia.
                                                       Gaps
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01-MAR-2004 (TrEMBLrel. 26, Created)
01-MAR-2004 (TrEMBLrel. 26, Last sequence update)
01-MAR-2004 (TrEMBLrel. 26, Last annotation update)
Glycine cleavage system protein H (Fragment).
Oryza sativa (Rice).
Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
Ehrhartoideae; Oryzeae; Oryza.
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                Score 17; DB 1; Length 9; Pred. No. 1.6e+06;
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                                                                                                                                                                                                                                                                    05-JUL-2004 (TrEMBLrel. 27, Created)
05-JUL-2004 (TrEMBLrel. 27, Last sequence update)
05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
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                                                       1; Mismatches
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Submitted (APR-1993) to the PIR data bank.
PIR; P80253; P80253.
Gramene; Q7M1V3; -.
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                                                                                                                                                                                                                                                                                                                               Ribosomal protein (Fragment).
              33.3%;
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                                                       3; Conservative
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                                                                                                                                                                                                                                                                                                                                                                    Alvesia rosmarinifolia.
Chloroplast.
            Query Match
Best Local Similarity
Matches 3; Conserv
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GLFRG 8
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1 GIVHL 5
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Chen W.M., Moulin L., Bontemps C., Vandamme P., Bena G.,
Bolyin-Masson C.;
"Symbiotic Nitrogen Fixation by beta-Proteobacter ia is widespread in
                             Gapa
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MEDLINE=80804539, PubMed=9392829, DOI=10.1016/S0196-9781(97)00188-5;
MEDLINE=80804539, PubMed=9392829, DOI=10.1016/S0196-9781(97)00188-5;
Duve H., Johnsen A.H., Maestro J.-L., Scott A.G., Winstanley D.,
Davey M., East P.D., Thorpe A.;
"Lepidopteran peptides of the allatostatin superfamily.";
Peptides 18:1301-1309(1997).
-1- SIMILARITY: Belongs to the allatostatin family.
Amidation; Direct protein sequencing; Neuropeptide.
Amidation; Direct protein sequencing; Neuropeptide.
ARGŪTENCE 7 AA; 873 MW; 672879CABB569350 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Sinorhizobium sp. BR816.
Bacteria; Proteobacteria; Alphaproteobacteria; Rhizobiales;
Rhizobiaceae; Sinorhizobium/Ensifer group; Sinorhizobium.
NCBI_TaxID=142626;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match 29.4%; Score 15; DB 1; Length 7; Best Local Similarity 66.7%; Pred. No. 1.6e+06; Matches 2; Conservative 1; Mismatches 0; Indels
                       Indels
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                                                                                                                                                                                                                                                                                                                                   05-JUL-2004 (TrEMBLrel. 27, Created)
05-JUL-2004 (TrEMBLrel. 27, Last sequence update)
05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
Pred. No. 1.6e+06;
3; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 31.4%; Score 16; DB 2; 1
75.0%; Pred. No. 1.6e+06;
tive 0; Mismatches 1;
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30-MAY-2000 (Rel. 39, Last sequence update)
05-JUL-2004 (Rel. 44, Last annotation update)
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EMBL; AJ300234; CAC82843.1; -.
NON TER 9 9
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                    2; Conservative
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Matches 3; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                               NodB protein (Fragment).
                                                                                                                                                                                                                                                                                 PRELIMINARY;
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Job time : 174 secs

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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Takahashi T., Furukawa Y., Muneoka Y., Matsushima O., Ikeda T., Fujita T., Minakata H., Nomoto K.; Fujita T., Minakata H., Nomoto K.; Isolation and characterization of four novel bloactive peptides from a polychaete annelid, Perinereis vancaurica."; Submitted (SEP-1955) to the EMBL/GenBank/DDBJ databases. SEQUENCE 8 AA; 989 MW; 954772CAA87B0B59 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Apre S.K., Uhlemann B., Schmid R., Altendorf K.;
Submitted (OCT-2001) to Swiss-Prot.
-1- FUNCTION: Apoprotein for the iron-sulfur centers FA and FB of the
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Gaps
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01-DEC-2001 (TrEMBLrel. 19, Last sequence update)
01-DEC-2003 (TrEMBLrel. 15, Last annotation update)
Photosystem I iron-sulfur center (Photosystem I subunit VII) (9 kDa polypeptide) (PSI-C) (Fragment).
Anabaena sp. (strain L31).
Bacteria, Cyanobatteria, Nostocales, Nostocaceae, Anabaena.
NCBL_TaxID=29412;
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GO; GO:0015979; P:photosynthesis; IEA.
InterPro; IPR001450; 4Fe45 FERREDOXin.
PROSITE; PS00198; 4Fe45 FERREDOXIN; PARTIAL.
4Fe-45; Iron-sulfur; MeTal-binding; Photosynthesis; Photosystem I.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  photosystem I complex.
--- SUBCELLULAR LOCATION: Cytoplasmic.
--- SIMILARITY: THE IRON-SULFUR CENTERS ARE SIMILAR TO THOSE OF BACTERIAL-TYPE 4FE-4S PERREDOXINS.
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01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
01-MAY-2000 (TrEMBLrel. 13, Last annotation update)
BIOACTIVE peptide P4-PUTATIVE ESOPHAGEAL NEUROREGULATOR.
Perinereis vancaurica.
BUKATYOTA; MELAZOA; Annelida; Polychaeta; Palpata; Aciculata; Phyllodocida; Nereididae; Perinereis.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match 29.4%; Score 15; DB 2; Length 8; Best Local Similarity 66.7%; Pred. No. 1.6e+06; Matches 2; Conservative 1; Mismatches 0; Indels
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Best Local Similarity 66.7
Matches 2; Conservative
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